



Whitelaw Compliance Group, LLC.

Examination of Compliance Standards for Opioid Manufacturers and Distributors

Prepared For	Prepared By
<p>UNITED STATES DISTRICT COURT FOR THE NORTHERN DISTRICT OF OHIO, EASTERN DIVISION</p> <p><i>IN RE NATIONAL PRESCRIPTION OPIATE LITIGATION</i></p> <p>Case No. 18-OP-45132 (N.D. Ohio) MDL No. 2804 Case No. 17-md-2804 Judge Dan Aaron Polster</p>	<p>Dr. Seth B. Whitelaw</p> <p>President & CEO Whitelaw Compliance Group, LLC.</p> <p>April 15, 2019</p>

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PART I: Qualifications, Scope & Methodology



1 Qualifications

For the past 30 years, I have worked in the life sciences industry as a food and drug attorney, compliance officer, compliance consultant and professor. In addition to my J.D., I have an LL.M. in Administrative Law and an S.J.D. in Health Law. Consequently, I have extensive experience working with and interpreting legislation, statutes, regulations and guidance documents.

Since 1993, I have designed, built, and run four separate corporate compliance programs for both pharmaceutical and medical device manufacturers (C.R. Bard, Inc., SmithKline Beecham Pharmaceuticals NA, GlaxoSmithKline R&D, Misonix, Inc.). I also teach monitoring and auditing to law students and working professionals, who are enrolled in Mitchell Hamline School of Law's Healthcare Compliance Certificate program.

As a consultant for Deloitte and now my own firm, I have assessed the effectiveness of numerous U.S. and international compliance programs and their ability to detect and prevent violations of the various legal, regulatory and industry standards that govern life science company operations. In addition to assessing or developing the full compliance program, I have assessed and helped develop controls in numerous discrete areas including, but not limited to:

- pharmaceutical sampling,
- payments to and services from healthcare professionals ("HCPs"),
- product diversion controls ("grey market"),
- laboratory controlled substances controls,
- promotional material claims and use,

- third-party qualification, contracting and use, and
- medical affairs unsolicited request systems.

As an in-house compliance officer, I have conducted many audits and internal investigations directed at uncovering specific misconduct by individuals at all levels of the organization. These investigations have involved sample diversion, product diversion, clinical trial fraud, bribery and corruption, theft and misuse of human biospecimens, and the falsification of domestic and international regulatory documents (submissions, reports, certifications, licenses, import/export documents, etc.).

None of the organizations reviewed in this report have employed me or engaged the services of me and my firm. For my services on this project, I am billing \$400 per hour. My compensation is not dependent on my testimony or on the outcome of this case. All my opinions offered in this report are offered to a reasonable degree of certainty. Also, I reserve the right to modify or supplement my conclusions as additional information becomes available to me, or as I perform further analyses.

2 Scope & Methodology

2.1 Scope

As an expert in the design, implementation, and operation of compliance programs in the life science industry, I was retained to examine, review and discuss:

1. The relevant standards surrounding the design, implementation, and operation of corporate and controlled substances compliance programs for the pharmaceutical industry.¹
2. The application of those standards to manufacturers and distributors² of controlled substances.
3. The effectiveness of the compliance programs for five distributors and one manufacturer of prescription opioid medicinal products based upon available documentation from 1996 to 2018 (“review period”).

¹ The term pharmaceutical industry is used to encompass both pharmaceutical manufacturers (“marketing defendants”) and the distributors of finished pharmaceutical products to physicians, hospitals, clinics and pharmacies (“distributor defendants”).

² Within the pharmaceutical supply chain from manufacturer to patient, pharmaceutical distributors occupy the mid-point of the chain. Thus, at the most basic level, distributors handle the logistics of getting medicinal products from the manufacturers to the local pharmacies (including hospitals and clinics) that dispense or fill the patient’s prescription obtained from a licensed prescriber (doctor, dentist, nurse practitioner, physician’s assistant, etc.).

2.2 Methodology

The manufacturers and distributors of opioids (listed as Schedule II or III controlled substances) reviewed in this report can be further categorized into groups by the type of business model. As a result, there are three different groups reviewed in this report.

- Group 1 (“G1”) distributors have a standard, “pure” distribution business model, which only involves distributing pharmaceutical products and providing other ancillary data and logistical services (not in the scope of this review). These distributors, McKesson, Cardinal Health and AmerisourceBergen, also are known as the “Big Three.”
- The Group 2 (“G2”) distributors have a standard business model that involves embedding distribution operations within a large, national pharmacy chain that supplied only its own pharmacies with opioid products. This group of distributors also utilize the G1 distributors to ensure an uninterrupted supply of opioids to their pharmacies or to handle Schedule II controlled substances. The G2 distributors examined are CVS and Walgreens.
- The Manufacturer Group produce the finished opioid products and typically sell in bulk quantities to the G1 distributors to supply retail pharmacy outlets. Mallinckrodt was sole member of this group.

Based on my experience and expertise outlined above, I can fairly evaluate the compliance controls employed by manufacturers and distributors and render opinions on whether they are aligned with regulatory requirements, expectations and leading industry practices, as well as whether they can be considered effective. My approach to this review utilized the same methodology I have used during the last 30 years when auditing or investigating compliance issues.

For all three groups in order to gain an understanding of each company’s corporate compliance and anti-diversion programs during the review period, I conducted a detailed examination of both publicly available statements and documents, and documents produced by the manufacturers and distributors in the course of this case. In the course of preparing this report, that information included, but was not limited to:

- company websites and press releases;
- government enforcement settlement documents, including inspection reports, Memoranda of Agreement;
- government correspondence to and from the company;
- company policies and procedures;
- organization charts;
- reports of compliance breaches and investigations;
- compliance training materials;
- committee reports and presentation materials;
- audit and other internal review reports; and
- third party consultant reports.

That information examined was then evaluated against the standards described in Part II of this report.

I also examined relevant data showing opioid shipments as well as suspicious orders reported to the DEA by the distributors and manufacturers during the review period. This data pertained not only to Summit and Cuyahoga

Counties, but also other jurisdictions as well such as West Virginia. Although Summit and Cuyahoga Counties are the primary focus of this report, these anti-diversion programs were national programs and not state or county specific. Therefore, I have reviewed and evaluated activity that also occurred outside of Summit and Cuyahoga Counties. This is the same approach taken by the House Energy and Commerce Committee in its 2018 report.³

Finally, I also consulted with James Rafalski, a retired DEA diversion investigator, who also is an expert in this case. I discussed with him how the DEA applies the Controlled Substances Act, the accompanying regulations and the Agency's guidance when inspecting the controlled substances anti-diversion efforts of a manufacturer or a distributor, including their suspicious order monitoring programs. We also discussed what the DEA generally considers to constitute an effective controlled substances compliance program for a prudent registrant.

PART II: Compliance Program Standards



3 Understanding the Context

This part of the report discusses the compliance standards that pertain to the marketing, sale, and distribution of prescription opioid products. Although the focus of this report is on prescription opioid products, and with good reason given the current public health crisis,⁴ most of the applicable compliance programs standards are not opioid specific. Likewise, these standards are publicly available and routinely accessed by compliance

³ See U.S. House Energy & Commerce Committee Report, *Red Flags and Warning Signs Ignored: Opioid Distribution and Enforcement Concerns in West Virginia*, 115th Cong., 9 (Dec. 19, 2018) (While focused on a narrow part of West Virginia, the report raises grave concerns about practices by the distributors and the DEA nationwide.) [“W.Va. Red Flags Report”].

⁴ See Discussion *infra* at Appendix A, Figure 1.

professionals in the pharmaceutical industry to evaluate and develop industry-specific corporate and controlled substances compliance programs.

3.1 General Overview of Compliance

Within the pharmaceutical industry, the term “compliance” is used to describe a vast array of functions and activities. For example, there is “Quality Compliance,” “Regulatory or FDA Compliance,” controlled substances compliance (a.k.a. “Suspicious Order Monitoring” or “SOM”) and others. However they are described, these functions are focused on integrating into a company’s business fabric, values and principles, as well as societal expectations expressed through laws, regulations, and industry standards of conduct.⁵ Therefore, compliance is not simply focused on legal and regulatory compliance to create an organizational framework to detect and prevent illegal or unethical conduct, but with establishing and promoting a corporate culture that manages risk, protects the company’s reputation, and above all strives to do no harm (*primum non nocere*). This is the true essence of “compliance.”

A primary function of the corporate compliance (a.k.a. “Big C Compliance”) program is oversight and coordination of the other internal compliance functions to minimize duplication, together with providing an organization’s Board of Directors and senior management a contextualized picture of the organization’s compliance posture at a given moment of time, which highlights areas where the organization’s behavior can improve.

This case concerns compliance standards for the marketing, sale, and distribution of prescription opioid products. While all prescription products carry some degree of risk, prescription opioid products, even when used for legitimate medicinal purposes, pose a special level of risk given their propensity to cause harm through addiction and the risk of diversion into the “black market” of illegal drugs.

This is evidenced by the fact that not only do prescription opioids require a prescription from a licensed medical professional, but they also have additional government-imposed controls surrounding the distribution and dispensing of these products. Therefore, it is expected that any company involved in the marketing, sale, or distribution of these products maintains a robust and effective compliance function in accordance with values, principles and societal expectations that strive to do no harm by ensuring these products are obtainable by legitimate patients while maintaining efforts to detect and prevent illegal diversion. As an ancillary benefit, such efforts can reduce the company’s exposure to legal and reputational risk helping the company fulfill its “contract” with shareholders to protect their investments and maintain confidence in the company.

This expectation to maintain a robust and effective compliance program is true even if there were no laws or regulations governing the marketing, sale, and distribution of prescription opioid products (e.g., the Controlled Substances Act). It also is an important “compliance” consideration that laws and regulations constitute the “floor” and not the “ceiling” of expected conduct. In other words, laws and regulations establish the bare minimum requirements that companies must abide by, but good companies, especially those that understand the

⁵ See, e.g. Brent Saunders, Our Social Contract with Patients, Allergan CEO Blog (Sep. 6, 2016), <https://www.allergan.com/news/ceo-blog/september-2016/our-social-contract-with-patients.aspx>. Brent Saunders, current CEO of Allergan Plc and a former Compliance Officer for Schering-Plough Corporation wrote “[t]he health care industry has had a long-standing unwritten social contract with patients, physicians, policy makers and the public at large.”

value of compliance, can and often do go farther.

The specific governing standards for what constitute effective compliance programs for the pharmaceutical industry are derived largely from four source categories. These are (1) state and federal statutes, plus any accompanying regulations, (2) government guidance documents, (3) industry enforcement settlements, and (4) voluntary industry standards including codes of conduct or ethics. Compliance professionals use these categories to develop compliance programs in the pharmaceutical industry that manage legal, regulatory and reputational risks effectively.

Finally, it is important to keep in mind that the origins of both controlled substances and corporate compliance programs predate the start of the report's review period (i.e., 1996).

3.2 The Interlocking Relationship between Suspicious Order Monitoring, Controlled Substances, and Corporate Compliance Programs

A suspicious order monitoring or SOM program is a subset of the wider universe of controls necessary for a distributor to meet its overall obligation to maintain "effective controls against diversion of particular controlled substances into other than legitimate medical, scientific, and industrial channels."⁶ As discussed in greater detail below, that wider universe of controlled substances diversion controls is itself a subset of the universe of controls a distributor needs to meet its ability to exercise due diligence to prevent and detect inappropriate and potentially criminal conduct and to promote otherwise an organizational culture that encourages ethical conduct (a.k.a. a corporate compliance program). The figure below illustrates that the relationship between SOM, a full controlled substances program and the enterprise-wide corporate compliance program resembles a set of Russian nesting dolls.

⁶ See 21 U.S.C. §§ 823 (b)(1). The Uniform Controlled Substances Act of 1994 states that "'diversion' means the transfer of a controlled substance from a lawful to an unlawful channel of distribution or use." See National Conference of Commissioners on Uniform State Laws, *Uniform Controlled Substances Act (1994)*, § 309(a) (Dec. 28, 1995) at http://www.uniformlaws.org/shared/docs/controlled%20substances/UCSA_final%2094%20with%2095amends.pdf.

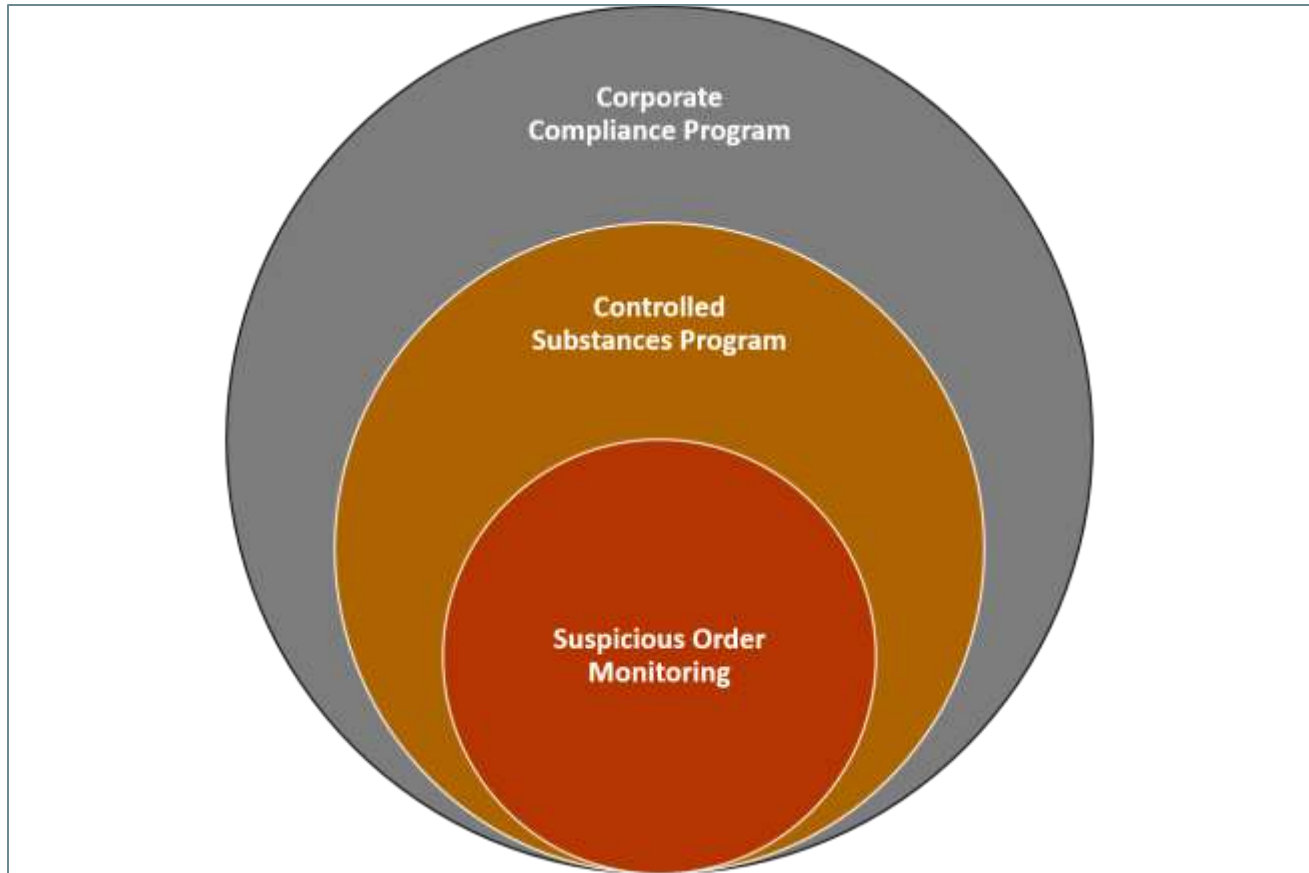


Figure 1: Relationship Between SOM, Controlled Substances & Corporate Compliance

As a result of this interlocking or “nested” arrangement, for a compliance program at any of the levels to be considered effective its basic building blocks must address the Seven, now Eight, Elements of an Effective Compliance Program.

4 Compliance Standards for Corporate Compliance Programs (1991 to the Present)

4.1 Federal Sentencing Guidelines for Organizations

In 1991, twenty years after passage of the CSA, the modern corporate compliance program was born with the publication of the first version of the Federal Sentencing Guidelines for Organizations (“FSGs”). Established by the U.S. Sentencing Commission (“Sentencing Commission”), the Guidelines are a “mechanical structure [that] determines an appropriate monetary fine through means of a mathematical formula: assigning a dollar figure to the seriousness of the offense and multiplying that number by a figure representing the culpability level of the

organization.”⁷ Consequently, “[t]he Guidelines’ drafters intend[ed] to influence corporate behavior – both before and after wrongdoing occurs – by providing various adjustments to the determination of the seriousness of the offense and of the organization’s culpability.”⁸

Applying a “carrot and stick approach,” the Sentencing Commission gave organizations an incentive to implement an effective compliance program. Therefore, the FSGs:

not only encourage corporations to exemplify “good corporate citizenship,” but [they] also provide a means to “rehabilitate” corporations that have engaged in criminal conduct⁹

According to the FSGs, “[t]he hallmark of an effective [compliance] program to prevent and detect violations of law is that the organization exercises due diligence in seeking to prevent and detect criminal conduct by its employees and other agents.”¹⁰ The Sentencing Commission, in a comment to the applications section, outlined seven criteria for establishing an effective compliance program. Commonly referred to as the “Seven Elements,” the FSGs required that for a compliance program to qualify as “effective” and receive mitigation credits:¹¹

1. The organization must have established compliance standards and procedures to be followed by its employees and other agents that are reasonably capable of reducing the prospect of criminal conduct;
2. Specific individual(s) within the high-level personnel of the organization must have been assigned overall responsibility to oversee compliance with such standards and procedures;
3. The organization must have used due care not to delegate substantial discretionary authority to individuals whom the organization knew or should have known through the exercise of due diligence, had a propensity to engage in illegal activities;
4. The organization must have taken steps to communicate its standards and procedures effectively to all employees and other agents, by requiring participation in training programs or by disseminating publications that explain in a practical manner what is required;
5. The organization must have taken reasonable steps to achieve compliance with its standards, by utilizing monitoring and auditing systems reasonably designed to detect criminal conduct by its employees and other agents and by having in place and publicizing a reporting system whereby employees and other agents could report criminal conduct by others within the organization without fear of retribution;

⁷ See Lawrence Finder and A. Michael Warnecke, *Overview of The Federal Sentencing Guidelines for Organizations and corporate Compliance Programs*, 1, ABA Criminal Justice Section (Apr. 12, 2005) at https://www.americanbar.org/content/dam/aba/publishing/criminal_justice_section_newsletter/crimjust_wcc_OVERVIEW_OF_THE_FEDERAL_SENTENCING_GUIDELINES_FOR_ORGANIZATIONS_AND_CORPORATE.authcheckdam.pdf.

⁸ *Id.*

⁹ See Diane Murphy, *The Federal Sentencing Guidelines for Organizations: A Decade of Promoting Compliance and Ethics*, 87 IOWA L. REV. 697, 703 (2002) (citations omitted)

¹⁰ See *id.* (Quoting from the U.S. Sentencing Guidelines Manual at ch. 8).

¹¹ See U.S. Sentencing Commission, *Guidelines Manual*, § 8A.1.2, comment. (n. 3k) (Nov. 1991).

6. The standards must have been consistently enforced through appropriate disciplinary mechanisms, including, as appropriate, the discipline of individuals responsible for the failure to detect an offense. The adequate discipline of individuals responsible for an offense is a necessary component of enforcement; however, the form of discipline that will be appropriate will be case specific; and
7. After an offense has been detected, the organization must have taken all reasonable steps to respond appropriately to the offense and to prevent further similar offenses -- including any necessary modifications to its program to prevent and detect violations of law.

These general elements outlined in the Sentencing Guidelines are not pharmaceutical-specific but rather apply to corporations across all industries. As the Sentencing Commission noted in its commentary, “[t]he precise actions necessary for an effective program to prevent and detect violations of law will depend upon a number of factors” including, but not limited to the size of the organization, the fact that via the nature of the business certain types of offenses are more likely to occur, and the organization’s prior history.¹² Therefore, it is incumbent upon each corporation to implement the elements in a way that effectively addresses and mitigates the risks in their specific industry and for their individual company.

From their origin in 1991 through 2010, the Seven Elements were not legally or regulatorily mandated. Nevertheless, after the case of *U.S. v. C.R. Bard, Inc.* in 1994,¹³ many of the larger pharmaceutical companies and other health care organizations began voluntarily implementing the Seven Elements with an understanding that the elements established the foundation for determining the worthiness of their compliance efforts and programs.

During this initial phase, the focus of industry activity and government enforcement actions was largely confined to establishing the role of the compliance officer and instituting the basic compliance framework outlined by the Federal Sentencing Guidelines.¹⁴ The baseline requirements of a compliance program in this era typically involved:

1. Hiring a compliance officer and establishing a compliance committee;
2. Developing written compliance standards and policies;
3. Implementing an employee training program;
4. Establishing a confidential disclosure program (e.g., hotline);
5. Restricting the employment of ineligible persons (e.g., pre-employment screening); and
6. For companies under a plea agreement, providing implementation and annual reports to OIG on the status of the entity’s compliance activities.¹⁵

¹² See *id.*

¹³ 848 F. Supp. 287 (D. Mass. 1994).

¹⁴ *Id.* Although *Bard* was an FDA enforcement action against a medical device company, the settlement, which required the company to develop and implement a compliance program, helped motivate the pharmaceutical industry to make corporate compliance a priority.

¹⁵ These baseline requirements were later expanded in 2001 with the TAP Pharmaceuticals Corporate Integrity Agreement. See generally, Corporate Integrity Agreement between DHHS OIG and TAP Pharmaceutical Products, Inc., https://www.oig.hhs.gov/fraud/cia/agreements/tap_pharmaceutical_products_92801.pdf (2001). The TAP CIA saw the introduction of the Compliance Committee and the independent review organization (“IRO”) to conduct annual reviews, as well as the

In 2004, the Federal Sentencing Commission significantly updated the Sentencing Guidelines. The corporate compliance program section was improved and expanded clearly signaling the importance of corporate compliance programs. Perhaps most importantly, the Sentencing Commission elevated the corporate compliance discussion from a comment to its own new chapter and section.¹⁶ The Commission also made three other major changes.

First, with the addition of “ethics” to the program name, the Sentencing Commission signaled that these programs have an expanded role beyond just detecting and preventing criminal conduct. As of 2004, an effective ethics and compliance program was intended to promote “an organizational culture that encourages ethical conduct and a commitment to compliance with the law.”¹⁷

Second, the Seven Elements were expanded to include an eighth element – risk assessment. Although listed explicitly for the first time, the risk assessment element was implied in the original 1991 Guidelines comment.¹⁸ With the 2004 changes, the section explicitly highlighted it stating:

In implementing subsection (b), the organization shall periodically assess the risk of criminal conduct and shall take appropriate steps to design, implement, or modify each requirement set forth in subsection (b) to reduce the risk of criminal conduct identified through this process.¹⁹

Third, the Sentencing Commission explicitly articulated that courts and judges could apply industry practice and standards in government regulations when concluding whether a compliance program was effective, and the failure to take governmental guidance and industry standards into account was viewed as a negative.

Specifically, the Commission wrote:

- (A) **In General.**—Each of the requirements set forth in this guideline shall be met by an organization; however, in determining what specific actions are necessary to meet those requirements, factors that shall be considered include: (i) applicable industry practice or the standards called for by any applicable governmental regulation; (ii) the size of the organization; and (iii) similar misconduct.
- (B) **Applicable Governmental Regulation and Industry Practice.** —An organization’s failure

requirement to make self-disclosures to the OIG of overpayments, investigations, legal proceedings, and other “reportable events” defined by the agreement. That settlement also introduced the conjoined concepts of the “covered person” and “certain covered persons” targeting various groups of employees for additional scrutiny and training. Now, instead of just one class of employees requiring training, the TAP CIA, and its progeny, required companies to identify those employees who constituted “covered persons or “certain covered persons” and establish and track training programs, as well as certifications, tailored specifically to each group.

¹⁶ See U.S. Sentencing Commission, *Guidelines Manual*, § 8B.2.1 (Nov. 2004) [“FSGs 2004”]. Section 8B.2.1 was amended in 2010, 2011 and 2013, however, those amendments were technical in nature and did not affect the overall requirements set out in that section. See U.S. Sentencing Commission, *Guidelines Manual*, Appendix C and Supplement to Appendix C (Nov. 2018) (Amendments 744, 758 and 778). Therefore, the 2004 Sentencing Guidelines contain the last major substantive update to the compliance program section.

¹⁷ See *id.* at § 8B.2.1(a)(2).

¹⁸ Although this concept was noted in the 1991 version, it was the very last sentence of the comment. See U.S. Sentencing Commission, *Guidelines Manual*, § 8A.1.2, comment. (n. 3k) (Nov. 1991).

¹⁹ *Id.* at § 8B.2.1(c).

to incorporate and follow applicable industry practice or the standards called for by any applicable governmental regulation weighs against a finding of an effective compliance and ethics program.²⁰

4.2 OIG Compliance Program Guidance

From 1998 to 2008, the Office of Inspector General (“OIG”) for Health and Human Services issued a series of compliance program guidance documents that pertained to a wide variety of healthcare organizations and companies including hospitals, home health agencies and clinical laboratories in 1998²¹, durable medical equipment, and hospices in 1999²², pharmaceutical manufacturers in 2003²³ and nursing facilities in 2008.²⁴ According to the OIG, “[t]he purpose of the compliance program guidance is to encourage the use of internal controls to efficiently monitor adherence to applicable statutes, regulations and program requirements.”²⁵ Each guidance followed a standard pattern of discussing the elements of an effective compliance program, as articulated by the Sentencing Guidelines, in the context of that particular industry segment. The compliance program guidance also represented the OIG’s position on what constituted leading practices at that time for that industry segment.

Although the OIG never established specific compliance program guidance for pharmaceutical distributors, a close reading of the guidance published in 2003 for pharmaceutical manufacturers provides many informative insights suitable for distributors as well. In fact, the OIG noted that the information contained in the Guidance might be useful to other groups beyond just pharmaceutical manufacturers:

²⁰ See FSGs 2004. at § 8B.2.1, comment. (n. 2) (emphasis added). In 2005, the U.S. Supreme Court in a complex opinion concluded that the Sentencing Guidelines violated a defendant’s Sixth Amendment right to a jury, but also found that courts could still use them, provided the court was able to tailor the final sentencing to address the specific facts of the case. See *Finder* at 2 (Discussing *United States v. Booker*, 123 S. Ct. 785 (2005)). The DOJ in response issued a memorandum to all federal prosecutors instructing them that they are required to use the Sentencing Guidelines and ranges in all but the extraordinary case. See Memorandum from James B. Comey, Dep. Atty. Gen. To All Fed. Prosecutors, *Department Policies and Procedures Concerning Sentencing* (Jan. 28, 2005), available at http://sentencing.typepad.com/sentencing_law_and_policy/files/dag_jan_28_comey_memo_on_booker.pdf. The net result is that despite *Booker*’s holding, the Sentencing Guidelines continue to define the elements of an effective ethics and compliance program by the courts, the regulators and the life sciences industry, and the passage of the Affordable Care Act in 2010 (see below) has eroded *Booker*’s relevance even further.

²¹ See Department of Health and Human Services, Office of Inspector General, OIG Compliance Program Guidance for Hospitals, 63 Fed. Reg. 8987 (Feb. 23, 1998); OIG Compliance Program Guidance for Home Health Agencies, 63 Fed. Reg. 42410 (Aug. 7, 1998); OIG Compliance Program Guidance for Clinical Laboratories, 63 Fed. Reg. 45076 (Aug. 24, 1998). All OIG Compliance Guidance documents are available at <https://www.oig.hhs.gov/compliance/compliance-guidance/index.asp>. To date, the OIG has published no specific compliance program guidance document for distributors.

²² See Department of Health and Human Services, Office of Inspector General, OIG Compliance Program Guidance for the Durable Medical Equipment, Prosthetics, Orthotics, and Supply Industry, 64 Fed. Reg. 36368 (Jul. 6, 1999); OIG Compliance Program Guidance for Hospices, 64 Fed. Reg. 54031 (Oct. 5, 1999).

²³ See Department of Health and Human Services, Office of Inspector General, OIG Compliance Program Guidance for Pharmaceutical Manufacturers, 68 Fed. Reg. 23731 (May 5, 2003) (“OIG Pharma Guidance”).

²⁴ See Department of Health and Human Services, Office of Inspector General, OIG Compliance Program Guidance for Home Health Agencies, 63 Fed. Reg. 42410 (Aug. 7, 1998).

²⁵ See OIG Pharma Guidance at 23731.

In addition, the compliance program elements and potential risk areas addressed in this compliance program guidance may also have application to manufacturers of other products that may be reimbursed by federal health care programs.²⁶

As experienced compliance professionals know, any compliance program guidance does not necessarily need to be written for the specific industry segment to contain pertinent insights on what constitutes effective compliance.

4.3 Affordable Care Act

Perhaps the most significant change for corporate compliance programs occurred with the passage of the Affordable Care Act (“ACA”) in 2010.²⁷ As noted previously, the standards detailing what constitutes the make-up of an effective compliance program have existed since 1991 and were widely adopted by most large pharmaceutical manufacturers and other prudent life sciences companies. They also were incorporated in various government guidance documents and settlement agreements. However, the passage of the ACA now made having a corporate compliance program a requirement to be eligible to participate in and receive reimbursement from federal health care programs.

Under ACA section 6401(a)(7) in order to participate in the Medicare program (e.g., receive reimbursement) after an implementation date determined by the Secretary of HHS:

a provider of medical or other *items or services or supplier* within a particular industry sector or category shall, as a condition of enrollment in the program under this title, title XIX, or title XXI, establish a compliance program that contains the core elements established under subparagraph (B) with respect to that provider or supplier and industry or category.²⁸

The same requirement also was applied to participants in state Medicaid programs, as well as the Children’s Health Insurance Program (“CHIP”).²⁹

Congress used the concept of “core elements” to tie the previous corporate compliance guidance and standards into this new requirement by stating:

The Secretary, in consultation with the Inspector General of the Department of Health and Human Services, shall establish core elements for a compliance program under subparagraph (A) for providers or suppliers within a particular industry or category.³⁰

Congress also was specific regarding the Secretary’s implementation determination that:

²⁶ See OIG Pharma Guidance at 23742, n.5.

²⁷ See Patient Protection and Affordable Care Act, Pub. L. No. 111-148, § 6401(a)(7), 124 Stat. 119, 689 (codified as amended at 42 U.S.C. § 1320a-7h) (amending Part A of title XI of the Social Security Act by adding section 1128G) (2010) [hereinafter cited as ACA]

²⁸ *Id.* at § 6401(a)(7)(A) (Emphasis added).

²⁹ *Id.* at § 6401(b)(5) and (c)(2).

³⁰ *Id.* at § 6401(a)(7)(B).

The Secretary shall determine the timeline for the establishment of the core elements under subparagraph (B) and the date of the implementation of subparagraph (A) for providers or suppliers within a particular industry or category. **The Secretary shall, in determining such date of implementation, consider the extent to which the adoption of compliance programs by a provider of medical or other items or services or supplier is widespread in a particular industry sector or with respect to a particular provider or supplier category.**³¹

4.4 DOJ & OIG Program Effectiveness Guidance

In 2017, both the OIG and DOJ published guidance on the elements that they consider when determining whether a corporate compliance program is effective.³² However, “[b]oth sets of guidance emphasize that they are not a ‘checklist to be applied wholesale to assess a compliance program’ but rather are lists of common elements to be considered when ‘making an individualized determination.’”³³ In addition, while there are similarities between the two guidance documents, there also some significant differences starting with the format. The DOJ guidance is formulated as questions to be considered, while the OIG document examines things to measure and how to accomplish those measurements.

5 Compliance Standards for Controlled Substances (1970 – the Present)

5.1 Controlled Substances Act

The origins of effective compliance programs for controlled substances (a.k.a. anti-diversion programs) are traceable to the enactment of both the Controlled Substances Act (“CSA”), which is the primary statute governing the manufacture and distribution of controlled substances, and the Drug Enforcement Administration’s (“DEA”) implementing regulations.³⁴ Originally enacted in 1970, the CSA established the classification system for controlled substances (Schedules I-V), as well as general controls that pertain to each

³¹ *Id.* at § 6401(a)(7)(C) (Emphasis added). As of the date of this report, the Secretary has not issued a formal determination of “core elements” under subparagraph (B) or the implementation date under subparagraph (C). However, given the existence of the Federal Sentencing Guidelines, the OIG Compliance Program Guidance in 2003 and the most recent OIG and DOJ guidance documents on program effectiveness issued in 2017, I believe the pragmatic compliance reading is that the “core elements” and timing requirements have been satisfied. Consequently, as of 2010, any pharmaceutical distributor, which receives federal healthcare dollars either directly or indirectly, must have an effective corporate compliance program that addresses the risks in a particular industry or industry category.

³² See U.S. Department of Justice, Criminal Division- Fraud Section, “Evaluation of Corporate Compliance Programs,” <https://www.justice.gov/criminal-fraud/page/file/937501/download>; see also HCCA-OIG Compliance Effectiveness Roundtable, *Measuring Compliance Program Effectiveness: A Resource Guide* (Mar. 27, 2017) (“On January 17, 2017 a group of compliance professionals and staff from the Department of Health and Human Services, Office of Inspector General (OIG) met to discuss ways to measure effectiveness of compliance programs.”), available at <https://oig.hhs.gov/compliance/101/files/HCCA-OIG-Resource-Guide.pdf>.

³³ See S. Foroughi and K. Wildoner, *Effectiveness, The Holy Grail of Compliance - Both the DOJ & OIG Weigh In*, 3.7 LIFE SCIENCE COMPLIANCE UPDATE 7, 14 (Jul. 2017) (citations omitted), available at <https://complianceupdate.policymed.com>.

³⁴ See 21 U.S.C. § 801 *et seq.*, see also 36 Fed. Reg. 7778 (Apr. 24, 1971) codified at 21 C.F.R. part 1301.

schedule.³⁵ Regardless of the Schedule level, the fact that a medicinal product is scheduled means that it has been determined that additional controls regarding the manufacture, distribution, dispensing and prescribing of that product are necessary to safe guard the public health.³⁶

Schedule II products are defined as drugs with a high potential for abuse, with use potentially leading to severe psychological or physical dependence.³⁷ Consequently, products in this category are considered the most dangerous products that can be lawfully prescribed by a medical professional. Schedule III products are defined as drugs with a potential for abuse that is less than the drugs in Schedules I and II, with use potentially leading to moderate to low physical dependence and high psychological dependence.³⁸ While these products are considered less dangerous than Schedule II drugs, nevertheless, they are potent medicinal products requiring the additional controls mandated by the CSA to prevent diversion and misuse.

As a baseline, the CSA requires that all major participants in the controlled substance supply chain (manufacturers, distributors, dispensers, and prescribers) be registered, thus creating the so-called “closed loop” system.³⁹ It further defines the basic controls expected of both manufacturers and distributors. A critical condition for granting, and maintaining, a manufacturer’s or distributor’s registration is the “maintenance of effective controls against diversion of particular controlled substances into other than legitimate medical, scientific, and industrial channels.”⁴⁰ The failure of any registrant to “refuse or negligently fail to make, keep, or furnish any record, report, notification, declaration, order or order form, statement, invoice, or information required” by the Act is a criminal offense.⁴¹

Although the CSA has been amended several times since 1970, this basic requirement to maintain effective diversion controls has remained untouched.⁴² Therefore, when the manufacturers and distributors developed the governing standards of conduct to detect and prevent diversion of prescription opioids, it was incumbent on

³⁵ See 21 U.S.C. §§ 812(b)(2), (b)(3) and (c). The CSA defines an opioid as “any drug or other substance having an addiction-forming or addiction-sustaining liability similar to morphine or being capable of conversion into a drug having such addiction-forming or addiction-sustaining liability.” See 21 U.S.C. § 802(18). For purposes of this review the focus is opioid products, which are classified as Schedule II or III controlled substances.

³⁶ See, e.g., U.S. DEPARTMENT OF JUSTICE, DRUG ENFORCEMENT AGENCY, 96-2 DIVERSION INVESTIGATOR’S MANUAL, § 5126 (Apr. 16, 1996), CAH_MDL_02203357.

³⁷ See 21 U.S.C. § 812(b)(2).

³⁸ See 21 U.S.C. § 812(b)(3).

³⁹ See 21 U.S.C. § 823.

⁴⁰ See 21 U.S.C. §§ 823 (a)(1) and (b)(1) (Governing manufacturers and distributors respectively). As a threshold matter, the CSA does not specifically define “diversion.” However, the language “into other than legitimate medical, scientific, and industrial channels” infers that if a controlled substance were moved into an illegitimate channel that constitutes “diversion.” According to the Uniform Controlled Substances of 1994, “‘diversion’ means the transfer of a controlled substance from a lawful to an unlawful channel of distribution or use.” See National Conference Of Commissioners on Uniform State Laws, *Uniform Controlled Substances Act (1994)*, § 309(a) (Dec. 28, 1995) at http://www.uniformlaws.org/shared/docs/controlled%20substances/UCSA_final%2094%20with%2095amends.pdf.

⁴¹ See 21 U.S.C. § 842(a)(5).

⁴² See, e.g., Pub. L. 91-513 available at <https://www.gpo.gov/fdsys/pkg/STATUTE-84/pdf/STATUTE-84-Pg1236.pdf> (Comprehensive Drug Abuse Prevention and Control Act of 1970).

each of them to consider the CSA's requirements as they developed and maintained an effective anti-diversion program.

5.2 DEA Controlled Substances Regulations

A year after passage of the CSA, the DEA in 1971 issued implementing regulations to clarify many of the CSA's important provisions including the registration and security controls for manufacturers, distributors, and dispensers of controlled substances.⁴³ A crucial component for controlled substances distributors was the security controls section outlining the physical security and other controls for non-practitioners.⁴⁴

Building from the original CSA provisions, the DEA's regulations required that all non-practitioner registrants (e.g., manufactures and distributors) develop and maintain:

effective controls and procedures to guard against theft and diversion of controlled substances. In order to determine whether a registrant has provided effective controls against diversion, the Administrator shall use the security requirements set forth in Secs. 1301.72-1301.76 as standards for the physical security controls and operating procedures necessary to prevent diversion.⁴⁵

The types of security controls that manufacturers and distributors must employ include:⁴⁶

- Making a good faith inquiry to determine if the person or entity receiving controlled substances is authorized to receive them;
- Maintaining a system to detect and disclose suspicious orders (a.k.a. Suspicious Order Monitoring or SOM);
- Notifying the DEA of thefts or significant losses; and
- Ensuring that any common carriers used in the supply chain have adequate security measures to prevent losses.

As laid out by the DEA regulations, a manufacturer's and distributor's SOM program must meet a relatively short list of requirements:

- There must be a system designed and operated to disclose suspicious orders of controlled substances.
- The distributor must inform the local DEA Field Office when the distributor discovers a suspicious order.
- At a minimum, orders are deemed suspicious if they are (a) of unusual size, (b) deviate substantially from a normal pattern, or (c) of unusual frequency.⁴⁷

⁴³ See 36 Fed. Reg. 7778 (Apr. 24, 1971) codified at 21 C.F.R. part 1301.

⁴⁴ See 21 C.F.R. §§ 1301.72 and 1301.74.

⁴⁵ *Id.* at § 1301.71(a).

⁴⁶ See *id.* at 1301.74.

⁴⁷ See 21 C.F.R. § 1301.74(b); see also *Masters Pharmaceuticals, Inc. v. DEA*, 15-1335 (D.C. Cir. 2017) (upholding DEA's interpretations of its regulations relative to defining a suspicious order and the timing of reporting).

Additionally, any manufacturer, which provides complimentary samples, must maintain appropriate controls for controlled substances in addition to the general controls for pharmaceutical marketing samples.⁴⁸

Also embedded within the DEA's regulations was the important concept that effective security controls are not static.⁴⁹ The regulations expressly contemplated that security controls should be adjusted (increased or decreased) to account for changing circumstances.⁵⁰ When determining whether a registrant is in substantial compliance with the security requirements, the DEA may apply a variety of factors, including but not limited to, "[t]he adequacy of the registrant's or applicant's system for monitoring the receipt, manufacture, distribution, and disposition of controlled substances in its operations."⁵¹ Therefore, as of 1971, both manufacturers and distributors were on notice that, at a minimum, they were expected to assess their controls periodically (e.g., undertake a risk assessment), as well as maintain a system to detect suspicious orders of controlled substances.

5.3 DEA Guidance on Controlled Substances

5.3.1 Controlled Substances Security Manual & Suspicious Order Task Force (1997 to 2004)

In 1991, the DEA published the Controlled Substances Security Manual as an informational guide to the CSA.⁵² The manual provided a more user-friendly outline of the CSA and its accompanying regulations. Later, in November 1997, the DEA announced the formation of the Suspicious Order Task Force.⁵³ The task force was "responsible for providing the Attorney General with recommendations, advice, and proposals for the establishment of such guidelines that will adequately define suspicious orders of listed chemicals."⁵⁴ It was comprised of 20 members including members from "relevant industry/trade associations and state and local law enforcement agencies."⁵⁵

⁴⁸ *Id.* The PDMA, which is administered by the U.S. Food and Drug Administration ("FDA") governs the distribution of pharmaceutical marketing samples. FDA's regulations for the most part mirror the DEA's requiring proof of to whom the samples were delivered, reporting of significant losses, investigating losses and suspected falsification, and providing timely notification of losses to the Agency. *See* 21 C.F.R. part 203. One difference between the two regulatory schemes is that the FDA specifically mandates the manufacturer maintain written policies and procedures governing how its sample accountability systems and processes operate. *See* 21 C.F.R. § 203.34. In 2010, the ACA added section 6004 which added yet another layer to the sampling of non-scheduled pharmaceuticals moving them closer to the "closed loop" DEA system. *See* Patient Protection and Affordable Care Act, Pub. L. No. 111-148, § 6004, 124 Stat. 119, 689 (codified as amended at 42 U.S.C. § 1320a-7h) (amending Part A of title XI of the Social Security Act by adding section 1128G) (2010).

⁴⁹ *See, e.g.,* Letter from W. Goggin to J.M. Gray (Oct. 17, 2008) ("diversion control is not a 'one size fits all' effort), WAGMDL00673706.

⁵⁰ *Id.* at § 1301.71(c).

⁵¹ *Id.* at § 1301.71(b)(14).

⁵² *See* U.S. DEP'T OF JUSTICE, DRUG ENFORCEMENT ADMINISTRATION, CONTROLLED SUBSTANCES SECURITY MANUAL (May 1991) available at http://www.cogan.com/documents/DEA_Controlled_Substances_Security_Manual.pdf.

⁵³ *See* 62 Fed. Reg. 61829 (Nov. 19, 1997).

⁵⁴ *Id.*

⁵⁵ *Id.*

5.3.2 The Chemical Handler's Manual

The DEA created the Chemical Handler's Manual in response to the enactment of the various chemical control laws, amending the original CSA, but also to provide general guidance on complying with the CSA.⁵⁶ Therefore, it contains relevant guidance on diversion controls and suspicious orders, including suspicious order identification criteria established by the Task Force.⁵⁷

The Manual also outlined "a voluntary formula for use by distributors to wholesale and retail levels."⁵⁸ The formula outlined involved setting threshold purchase levels based on the last twelve months purchases by the same customer type from the same distribution center (e.g., the customer group).⁵⁹ That amount is divided by the total number of customer months (months in which purchases are above zero) and multiplied by a factor to determine the maximum amount a customer may purchase.⁶⁰ According to the Manual, the "[f]actor equals 3 for C-II and C-III Controlled Substances **containing List I Chemicals** and 8 for C-III-IV-V Controlled Substances and non-Controlled OTC products **containing List I chemical items**."⁶¹

While the manufacturers and distributors here utilized the Factor of 3 for setting thresholds for opioid products, the factor was based only on Schedule II and III controlled substances containing List 1 Chemicals.⁶² A plain reading of Appendix E-3 is that if a Schedule II or III controlled substance does not contain a List 1 chemical, the factor is not applicable. Therefore, for opioid products not containing a List 1 chemical, that factor is not applicable. However, regardless of whether using the factor is applicable or not, the DEA manual does not indicate how a level that is 300% above the base threshold is the appropriate multiplier to use.

Independent of the type of products the Chemical Handler's Manual applies to, it is clear that the Manual does not support the practice of shipping suspicious orders after they are reported. To this point, the Chemical Handler's Manual states "when a regulated person suspects that an order may be intended for illicit purposes, good practice requires that every reasonable effort be made to resolve those suspicious. In addition to making the required reports, the transaction should not be completed until the customer is able to eliminate the suspicions. The distributor may have to forego some transactions."⁶³

⁵⁶ See U.S. DEP'T OF JUSTICE, DRUG ENFORCEMENT ADMINISTRATION CHEMICAL HANDLER'S MANUAL, (Jan. 2004) at <https://www.justice.gov/sites/default/files/open/legacy/2014/05/09/2004-chemical-handlers-manual.pdf>. ["Chemical Handlers Manual, 2004 Edition"].

⁵⁷ *Id.* at 37 (Appendix E).

⁵⁸ *Id.* at 41 (Appendix E-3).

⁵⁹ *Id.*

⁶⁰ *Id.*

⁶¹ *Id.* (emphasis added).

⁶² The Manual states that a "*List I chemical* is a chemical that, in addition to legitimate uses, is used in manufacturing a controlled substance in violation of the CSA and is designated a List I chemical by the DEA Administrator or Congress. Chemicals in List I generally are precursors and have been determined by DEA to require a greater level of control than other listed chemicals." See *id.* at 8 (emphasis added).

⁶³ See Chemical Handler's Manual, 2004 Edition, at 19.

5.3.3 The DEA Industry Initiative

“Recognizing that wholesale distributors played a key role in the pharmaceutical supply chain, the DEA launched an industry-specific anti-diversion initiative in 2005, called the “Distributor Initiative Program.”⁶⁴ According to the DEA, the goal of the initiative was to “educate registrants on maintaining effective controls against diversion, and monitoring for and reporting suspicious orders.”⁶⁵ Initially, the DEA focused the program on educating drug distributors who were supplying controlled substances to rogue Internet pharmacies and to diverting pain clinics and pharmacies.⁶⁶ Through the program, the DEA “educates distributors about their obligations under the CSA, as well as provides registrants with current trends and ‘red flags’ that might indicate that an order is suspicious.”⁶⁷ McKesson, Cardinal Health, and Amerisource Bergen all attended sessions with the DEA.⁶⁸ The materials used in each meeting were almost identical.⁶⁹

During those meetings, the DEA told the participants that:

1. Reporting a suspicious order to the DEA does not relieve the distributor of its responsibility to maintain effective anti-diversion controls.
2. The DEA cannot tell distributors if an order is legitimate or not.
3. Distributors, therefore, must determine which orders are suspicious and decide whether to proceed with the sale.
4. If distributors know or suspect that controlled substances are being dispensed outside the course of professional practice shipments to those customers must stop immediately.
5. The DEA may revoke a distributor’s registration under public interest grounds.⁷⁰

Although couched in terms of distributors, because the requirements for manufacturers are the same, the DEA’s statements as part of this initiative would apply to them too.

⁶⁴ See MEMORANDUM FROM COMMITTEE MAJORITY STAFF, H.R. COMM. ON ENERGY AND COMMERCE, SUBCMTE. ON OVERSIGHT AND INVESTIGATIONS, HEARING ENTITLED “COMBATING THE OPIOID EPIDEMIC: EXAMINING CONCERNS ABOUT DISTRIBUTION AND DIVERSION,” 5, (May 4, 2018), <https://docs.house.gov/meetings/IF/IF02/20180508/108260/HHRG-115-IF02-20180508-SD002.pdf>.

⁶⁵ *Id.* (quoting from *Improving Predictability and Transparency in DEA and FDA Regulation: Hearing Before H. Comm on Energy & Commerce, Subcomm. on Health*, 113th Cong., (2014) (statement of Joseph T. Rannazzisi, Deputy Assistant Adm’r, Office of Diversion Control, U.S. Drug Enforcement Admin.)).

⁶⁶ *See id.*

⁶⁷ *See id.* (quoting from *Improving Predictability and Transparency in DEA and FDA Regulation: Hearing Before H. Comm on Energy & Commerce, Subcomm. on Health*, 113th Cong., (2014) (statement of Joseph T. Rannazzisi, Deputy Assistant Adm’r, Office of Diversion Control, U.S. Drug Enforcement Admin.)).

⁶⁸ See Memorandum to J. Rannazzisi from M. Mapes, Internet Presentation with McKesson Corp. on September 1, 2005 (Oct. 20, 2005), MCKMDL00496859; Presentation by M. Mapes and K. Wright to Cardinal Health, *Internet Pharmacy Data*, (Aug. 22, 2005), CAH_MDL2804_01457737; Presentation by M. Mapes and K. Wright to AmerisourceBergen, *Internet Pharmacy Data*, (Aug. 10, 2005), ABDCMDL00315887.

⁶⁹ *Id.*

⁷⁰ See Presentation by Mapes and Wright to AmerisourceBergen at ABDCMDL00315893-94, and ABDCMDL00315899.

5.3.4 DEA Letters to All Registrants (a.k.a. The Rannazzisi Letters) (2006 to 2012)

In 2006, 2007 and again in 2012, Joseph Rannazzisi, Deputy Assistant Administrator of the Office of Diversion Control also issued a series of guidance letters.⁷¹ Known collectively as the Rannazzisi letters, they were sent to all registered manufacturers and distributors reminding them of their obligations under the CSA to prevent diversion and detect suspicious orders.⁷² Beyond the general reminders and disclaimer that the DEA does not endorse a particular system or sets of controls, each letter focused on a particular implementation topic, providing DEA's current thinking about what was or was not effective.

The initial letter in September 2006 focused on a registrant's basic obligations noting that the suspicious order monitoring "requirement is in addition to, and not in lieu of, the general requirement . . . that a distributor maintains effective controls against diversion."⁷³ The DEA also provided a list of factors that could signal possible diversion.⁷⁴

The focus of the February and December 2007 letters again was suspicious order monitoring. While the February 2007 letter's content was almost identical to the September 2006 letter, the December 2007 letter focused on what constituted timely reporting.⁷⁵ The December letter also cautioned registrants about placing too much reliance on rigid formulas to detect diversion, as well as the need to conduct meaningful investigations of suspicious orders.⁷⁶

The June 2012 letter continued the discussions started in December 2007 and once more focused on suspicious order monitoring. This time the DEA expressed concerns over registrants' not making timely reports to the DEA Field Offices as the regulations require. However, the DEA commented that merely reporting suspicious orders was not enough noting:

Registrants who routinely report suspicious orders yet fill these orders without first ascertaining that the order will not be diverted into other than legitimate medical, scientific, or industrial channels, are failing to maintain effective controls against diversion.⁷⁷

Thus, the DEA reiterated its expectation that registrants needed to conduct meaningful due diligence before

⁷¹ See Letters from J. Rannazzisi to All Registrants (Sep. 27, 2006, Feb. 7, 2007, Dec. 27, 2007 and Jun. 12, 2012) ["DEA (date) Letter(s)"].

⁷² *Id.*

⁷³ See DEA 9/27/2006 Letter at 2.

⁷⁴ See *Id.* at 3 (Listing circumstances that might be indicative of diversion).

⁷⁵ See DEA 12/27/2007 Letter; see also Letter from G.Thomas Gitchel to R.J. Streck (Apr. 27, 1984) ("any automated data processing system may provide the means and mechanism for compliance when the data is carefully reviewed and monitored by the wholesaler."), CAH_MDL2804_01465723. Mr. Gitchel was the DEA's Acting Chief of the Diversion Operations Section at that time.

⁷⁶ *Id.*

⁷⁷ See DEA 6/12/2012 Letter at 2; see also HDMA *Position Statement and Industry Compliance Guidelines: Report Suspicious Orders and Preventing Diversion of Controlled Substances* (2008), WAGMDL00673706-WAGMDL00673722.

shipping potentially suspicious orders.⁷⁸

5.3.5 Masters Pharmaceutical Case

While the case revoking the DEA registration for Masters Pharmaceuticals, Inc. ultimately came before the D.C. Circuit,⁷⁹ the opinion of DEA's Acting Administrator Chuck Rosenberg provides specific guidance on the determination of exactly when an order of unusual size, frequency or pattern "is discovered" as "suspicious."⁸⁰ This determination is particularly important because if a suspicious order "is discovered," the manufacturer or distributor should not ship the order to the customer.⁸¹ Thus, as discussed throughout this report, distributors and manufacturers go to extraordinary lengths to avoid "discovering" a suspicious order.

The regulations do not expressly define what is meant by "when discovered," and as a result, manufacturers and distributors use various euphemisms, such as "orders of interest" or like terms not found in the regulation as an attempt to avoid triggering the reporting requirement. However, the general principles of statutory construction hold that words not defined by a statute or regulation should be given their "plain meaning" as derived from the dictionary.⁸² Consequently, when a manual or automated threshold system "discloses" the excessive/suspicious order that constitutes "when discovered" triggering the reporting requirement.

According to Mr Rosenberg's opinion "[s]uspicion as to the existence of a circumstance (i.e., that a customer is engaged in diversion) is simply a far lower standard of proof than whether it is 'likely' that the circumstance exists ... [and] does not even rise to the level of probable cause."⁸³ Thus, he concluded that "an order has been discovered to be suspicious and the regulation has been violated where the registrant has obtained information that an order is suspicious but then chooses to ignore that information and fails to report the order."⁸⁴

⁷⁸ *Id.* In same vein as Rannazzisi letters, James Arnold, Unit Chief, Regulatory Unit, DEA HQ, in June 2013, spoke about diversion controls at a conference for manufacturers, importers and exporters hosted by the DEA. *See* Presentation by James Arnold, *Effective Controls Against Diversion*, Manufacturer/Importer/Exporter Conference, (Jun. 2013) available at https://www.deaiversion.usdoj.gov/mtgs/man_imp_exp/conf_2013/. While largely a recap of the statute and regulations, Mr. Arnold made several important points during his talk. First, he stressed that the responsibility for identify suspicious orders is the registrant's, but once identified as suspicious, the order must not be shipped. *Id.* at slide 41. Second, Mr. Arnold noted registrants must know their customers and have determined if there are any "red flags" to doing business with them. These "red flags" can include any number of factors including, but limited to, the customer's location, news reports, etc. *Id.* at slides 42 to 53.

⁷⁹ *See Masters Pharmaceutical, Inc. v. DEA*, No. 15-1335, (D.C. Cir. 2017). The D.C. Circuit's opinion is relevant because the Court affirmed the positions taken by Acting Administrator Rosenberg.

⁸⁰ *See* 80 Fed. Reg. 55418 (Sept. 15, 2015).

⁸¹ *See* Letter from J. Rannazzisi to All Registrants (Jun. 12, 2012); *see also* HDMA, Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances, 11 (2008) (blocking "orders of interest"), WAGMDL00673706.

⁸² *See, e.g., Morissette v. United States*, 342 U.S. 246, 263 (1952); *FDIC v. Meyer*, 510 U.S. 471, 476 (1994) (In the absence of a statutory definition, "we construe a statutory term in accordance with its ordinary or natural meaning."); *see also* LARRY M. EIG, CONG. RESEARCH SERVICE, 97-589, STATUTORY INTERPRETATION: GENERAL PRINCIPLES AND RECENT TRENDS, 5-8 (2014).

⁸³ *See* 80 Fed. Reg. at 55478.

⁸⁴ *Id.*

With regards to the “when discovered” provision, Mr. Rosenberg concluded the provision is intended “to prevent manufacturers and distributors from simply filing “daily, weekly, or monthly” suspicious order reports” because “periodic reports delay the reporting of suspicious orders ... meaning that DEA cannot act quickly when necessary.”⁸⁵ However, he concluded that the purpose of the language is “to impose a time period for ‘informing’ the Agency about a specific suspicious order.”⁸⁶

Consequently, when a manual or automated threshold system “discloses” the excessive/suspicious order that constitutes “when discovered” and triggers the reporting requirement. However, it is reasonable to permit a brief investigatory period to avoid the submission of reports that have been flagged by the system, but clearly are not suspicious as determined through verifiable and documented means. Therefore, based on the guidance provided by Acting Administrator Rosenberg’s conclusions in the *Masters* case, it is my opinion that this investigatory period is less than a week. To permit a longer investigative period would only increase the likelihood that the DEA will be provided stale information if the order is ultimately reported as suspicious, which would run counter to the Agency’s ability to properly investigate the order. It is also clear that the registrant must not ship the order until it is determined not to be suspicious and if the registrant cannot make a determination within the investigatory period, the order must be reported to the DEA and canceled.

5.4 Industry Guidance

In 1987, the National Wholesale Druggists’ Association (“NWDA”) developed, with input from the DEA, a suspicious order monitoring program.⁸⁷ The NWDA program or system provided, among other things, that “[s]ingle orders of unusual size or deviation must be reported [to the DEA] immediately ... [t]he submission of a monthly printout of after-the-fact sales will not relieve a registrant from the responsibility of reporting those single excessive or suspicious orders. DEA has interpreted ‘orders’ to mean **prior to shipment**.”⁸⁸

Building on the guidance provided by the DEA, Healthcare Distribution Management Association (“HDMA”), in 2008, developed voluntary industry guidelines to provide clear direction on how to develop a compliant anti-diversion program.⁸⁹ These general guidelines, which must be adapted by each individual distributor, cover the critical anti-diversion topics including:

- Know Your Customer Due Diligence;
- Monitoring for Suspicious Orders;

⁸⁵ *Id.*

⁸⁶ *Id.*

⁸⁷ See NWDA “Suspicious Drug Order” Monitoring Program, THE PINK SHEET (May 11, 1987), <https://pink.pharmaintelligence.informa.com/PS011879/NWDA-SUSPICIOUS-DRUG-ORDER-MONITORING-PROGRAM>. The National Wholesale Druggists’ Association became the Healthcare Distribution Management Association (“HDMA”) in 2000 and in 2016 HDMA became the Healthcare Distribution Alliance (“HDA”). See HDA, *History*, <https://www.hda.org/about/hda-history> (last accessed Feb. 21, 2019).

⁸⁸ See Nat’l Wholesale Druggists’ Ass’n, NWDA Suspicious Order Monitoring System, 7 (Jun. 21, 1993) (emphasis added), CAH_MDL2804_01465723.

⁸⁹ See HDMA, Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances, 13 (2008), WAGMDL00673706.

- Suspend/Stop an Order of Interest Shipment;
- Investigation of Orders of Interest;
- File Suspicious Order Reports with DEA;
- Employees, Training and Standard Operating Procedures; and
- Additional Recommendations.⁹⁰

While the Compliance Guidelines incorporate the provisions found in the CSA, the DEA regulations, and the various guidance documents from DEA, they also add concepts not found in those documents. For example, it is in the HDMA guidelines that the term “orders of interest” appears. As defined by HDMA, “orders of interest” are “orders that warrant follow-up inquiry to determine whether they are suspicious.”⁹¹ Furthermore, it appears that the letter from Wendy Goggin, Chief Counsel for DEA, commending HDMA’s efforts, is where the industry gets the mistaken belief that DEA “endorsed” the Compliance Guidelines, including the “orders of interest” concept.⁹² Also, HDMA in the Industry Compliance Guidelines counseled, “[d]istributors are strongly encouraged to regard timeliness of reporting to DEA as a critical component in meeting the requirement to report ‘when discovered.’”⁹³

⁹⁰ *Id.* at 3.

⁹¹ *Id.* at 8.

⁹² *See*, Letter from W. Goggin to J.M. Gray (Oct. 17, 2008) (“diversion control is not a ‘one size fits all’ effort), WAGMDL00673706.

⁹³ *See* HDMA, Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances, 13 (2008) (emphasis added), WAGMDL00673706.

PART III: Defining What Good Looks Like



6 Applying the Standards

As discussed in Part II, by the mid-1990s, the concept of “what good looks like” was established both in the context of corporate and controlled substances (a.k.a. anti-diversion) compliance. From that point forward it was clear that companies in the pharmaceutical industry, including manufacturers and distributors of opioid products, could develop effective internal controls to achieve the objectives to prevent and detect criminal conduct by an organization’s employees and agents working on behalf of the organization and to guard against theft and diversion of controlled substances.⁹⁴

In the U.S., the basic regulatory construct for pharmaceuticals, regardless of the agency, is to provide the industry with “what” is expected, but not dictate “how” those expectations are achieved. The “how” is left to the individual organizations to determine the best methods to comply. This approach is true in the case of the OIG, DEA, and even the FDA.⁹⁵

⁹⁴ See Appendix B, Figures 2 and 3 for diagrams outlining a controlled substances compliance program (a.k.a. anti-diversion program) and a corporate compliance program.

⁹⁵ See, e.g., U.S. Sentencing Commission, *Guidelines Manual*, § 8A.1.2, comment. (n. 3k) (Nov. 1991) [“FSGs 1991”]; J. Rannazzisi letters to All Registrants (Sep. 27, 2006, Feb. 7, 2007, Dec. 27, 2007 and Jun. 12, 2012) (These letters were not McKesson specific but sent to all DEA registrants), MCKMDL00478906, MCKMDL00615308, MCKMDL00478910, MCKMDL00449807 [“DEA (date) Letter”]; U.S. Food and Drug Admin., Center for Drug Evaluation and Research, *Facts About the Current Good Manufacturing Practices (CGMPs)*, <https://www.fda.gov/Drugs/DevelopmentApprovalProcess/Manufacturing/ucm169105.htm>, (page last updated Jun. 25, 2018) (last accessed Dec. 8, 2018) (“The CGMP requirements were established to be flexible in order to allow each manufacturer to decide individually how to best implement the necessary controls by using scientifically sound design, processing methods, and testing procedures.”).

6.1 General Principles

6.1.1 Corporate Compliance Programs

For any compliance program to be considered effective its basic building blocks must address the Seven, now Eight Elements of an Effective Compliance Program. These elements, whether for enterprise-wide or for controlled substances, are:

1. Organization and Resources
2. Due Diligence
3. Written Standards
4. Training & Communication
5. Monitoring, Auditing & Investigations
6. Corrective Actions
7. Enforcement (i.e., Discipline or other consequences for violating the standards)
8. Periodic Risk Assessment

While the eight elements provide a generally accepted and cohesive framework to assess compliance effectiveness, there is overlap between them, and therefore separating specific compliance activities by element is something of an esoteric exercise.

For purposes of simplicity and consistency when looking holistically across the entities assessed in my report, I grouped the eight elements listed above as follows:

Table 5.1-1 – Grouping the Eight Elements

Category	Elements of an Effective Compliance Program
Company Commitment	1. Organization and Resources (including company culture)
Program Core	3. Written Standards 4. Training & Communication 5. Monitoring, Auditing & Investigations 6. Corrective Actions 8. Periodic Risk Assessments
Accountability	2. Due Diligence (i.e., avoiding bad actors) 7. Enforcement (i.e., Discipline or other consequences for violating the standards)

Furthermore, although each type of compliance program has a specific focus (general enterprise-wide compliance versus controlled substances distribution compliance versus suspicious order monitoring), the

detailed standards applicable to all three types of compliance programs discussed here should be read together, as they reinforce and build-off each other.

Since the mid-1990's little has changed in the fundamentals in either the corporate compliance or controlled substances spheres, rather Main Justice, the DEA, and the OIG have become increasingly more pointed in reminding the pharmaceutical industry of what its statutory and regulatory obligations are with respect to corporate and controlled substances compliance. Even the introduction of new technology (e.g., ARCOS) has done little to change the fundamental compliance dynamic operating since 1995.

For example, in the context of monitoring compliance, technology arguably increases the amount of information that can be sorted, filtered and rapidly transmitted, but even today it merely provides an output of outliers and anomalies. Therefore, corporate and controlled substances compliance programs must still rely on experienced human resources, with intelligence and common sense, to review and understand the context surrounding each outlier or anomaly and then to apply the correct, balanced solution. Thus, in the end, good compliance comes down to experienced people making good choices.

It also comes down to the need for "objective evidence" to demonstrate that required compliance obligations including effectiveness, have been met. Thus, written documentation is the bedrock of demonstrating or "proving" that an organization's claims of effectiveness (or lack of thereof) are real. For example, as the Public Company Accounting Oversight Board ("PCAOB") points out in the context of audits:

Inadequate audit documentation diminishes audit quality on many levels. First, if audit documentation does not exist for a particular procedure or conclusion related to a significant matter, its absence casts doubt as to whether the necessary work was done.⁹⁶

The same applies to compliance efforts. Placing the PCAOB's comments about audit documentation into a compliance context:

Inadequate **compliance** documentation diminishes **compliance** quality on many levels. First, if **compliance** documentation does not exist for a particular procedure or conclusion related to a significant matter, its absence casts doubt as to whether the necessary work was done.

This is the same point made by the House Energy and Commerce Committee report.⁹⁷

Thus, if there is no documentation showing what is claimed was accomplished, the reasonable presumption is that it was not accomplished. Consequently, underlying all the applicable standards is the presumed need for good, written documentation to substantiate the existence and proper operation of compliance controls.

⁹⁶ See PCAOB, *Audit Documentation and Amendment to Interim Auditing Standards*, PCAOB Release No. 2004-006 (Jun. 9, 2004) (Announcing adoption of Audit Standard No. 3 on audit documentation). Reference to the PCAOB is appropriate in this context because most of the distributors reviewed are publicly traded entities and thus must arrange for independent audits of their financial statements. For those entities that are privately-held, there remains a basic fiduciary duty to the owners and company directors that also necessitates good documentation exists.

⁹⁷ See WVA Red Flags Report at 124-125, 130 and 319 (repeatedly commenting on the lack of due diligence documentation by distributors).

6.1.2 Suspicious Order Monitoring Programs

Overall, a distributor's SOM program must meet a relatively short list of requirements:

1. There must be a system designed and operated to disclose suspicious orders of controlled substances.
2. The distributor must inform the local DEA Field Office when the distributor discovers a suspicious order.
3. At a minimum, orders are deemed suspicious if they are (a) of unusual size, (b) deviate substantially from a normal pattern, or (c) of unusual frequency.
4. Suspicious orders must be held and not shipped until it is determined that the order likely will not be diverted.⁹⁸

As a threshold matter, the distributor or manufacturer must determine if the controlled substances customer is properly licensed to possess the controlled substance.⁹⁹ Both must also take steps to "know the customer," in other words, they need:

to take reasonable measures to verify the identity of their customers, understand the normal and expected transactions typically conducted by those customers, and, consequently, detect those transactions that are suspicious in nature.¹⁰⁰

As noted throughout this report, the "Know Your Customer" or KYC concept is critical to having a successful SOM program. To be effective, distributors and manufacturers must build and maintain profiles of their customers that are more specific than segregating those customers into various classes of trade. For example, knowing a pharmacy's product mix of controlled versus non-controlled prescriptions together with local data such as the number of residents, age as a percentage of residents, number and type of physicians and healthcare facilities, etc., are all important pieces of information that can make up a "Know Your Customer" profile. As the DEA makes clear, the Know Your Customer requirement is the basis for determining whether a customer's purchases are to be considered legitimate or diversionary. However, it also is important to remember that knowing one's customer and making determinations of whether orders are suspicious or legitimate is not simply a scientific endeavor (e.g., just using thresholds and algorithms), but also is an art requiring training, experience, innate skepticism, and common sense.

However, detecting and subsequently reporting suspicious orders are just a part of the overall set of controls a distributor and manufacturer needs to employ to prevent diversion. If "diversion" is taken to mean moving controlled substances into illegitimate "medical, scientific, [or] industrial channels"¹⁰¹ or if it is taken to mean "the transfer of a controlled substance from a lawful to an unlawful channel of distribution or use,"¹⁰² then to

⁹⁸ See 21 C.F.R. § 1301.74(b); see also *Masters Pharmaceuticals, Inc. v. DEA*, 15-1335 (D.C. Cir. 2017) (upholding DEA's interpretations of its regulations relative to defining a suspicious order and the timing of reporting).

⁹⁹ See 21 C.F.R. § 1301.74(a).

¹⁰⁰ See U.S. Dep't. of Justice, Drug Enforcement Administration *Chemical Handler's Manual*, 21 (2013) at https://www.dea diversion.usdoj.gov/pubs/manuals/chem/chem_manual.pdf.

¹⁰¹ See 21 U.S.C. §§ 823 (b)(1).

¹⁰² See Nat'l Conf. of Commissioners on Uniform State Laws, *Uniform Controlled Substances Act (1994)*, § 309(a) (Dec. 28, 1995) at http://www.uniformlaws.org/shared/docs/controlled%20substances/UCSA_final%2094%20with%2095amends.pdf.

prevent potential diversion, one needs to ensure that suspicious orders are not shipped until an appropriate investigation concludes that the risks of diversion occurring are not present.¹⁰³

Taking it one step further, since maintaining effective controls against diversion is just part of a manufacturer and distributor's overall responsibility to exercise due diligence to prevent and detect criminal conduct, the controlled substances program and suspicious order monitoring system need to have controls, including but not limited to, conducting periodic risk assessments and undertaking appropriate corrective actions that flow from the company's compliance standards. Otherwise, that distributor or manufacturer cannot contend that it has maintained effective controls against diversion or that its corporate compliance efforts are effective.

6.2 Compliance Culture, Organization & Resources

The 2004 Federal Sentencing Guidelines ("FSGs") mandate that for an ethics and compliance program to be considered effective, it must promote "an organizational culture that encourages ethical conduct and a commitment to compliance with the law."¹⁰⁴ As the OIG explained the year before in its 2003 Compliance Program Guidance, promoting and encouraging a commitment to ethics means:

for a compliance program to be effective, it must have the **support and commitment of senior management** and the company's governing body. In turn, the corporate leadership should strive to **foster a culture** that promotes the prevention, detection, and resolution of instances of problems.¹⁰⁵

For any compliance program to be successful, it must have adequate resources and the authority to achieve real compliance, and not just be delegated the responsibility for compliance.¹⁰⁶ In other words, responsibility without actual authority and appropriate resources is meaningless. Therefore, the culture of an organization, as well as the structure and resources are important elements of an effective compliance program.

Under the 2004 FSGs, in addition to requiring that high-level personnel in the organization be assigned responsibility for a compliance program, the Guidelines mandate:

- The organization's governing authority shall be knowledgeable about the content and operation of the compliance and ethics program and shall exercise reasonable oversight with respect to the implementation and effectiveness of the compliance and ethics program [and]

¹⁰³ See Letter from Wendy Goggin to John Gray (Oct. 17, 2008) (discussing HDMA's voluntary industry guidelines, "Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances.") WAGMDL00673706.

¹⁰⁴ See FSGs 2004 at § 8B.2.1(a)(2).

¹⁰⁵ See Dept. of Health and Human Services, Office of Inspector General, OIG Compliance Program Guidance for Pharmaceutical Manufacturers, 68 Fed. Reg. 23731 (May 5, 2003) (emphasis added) ["OIG Pharma Guidance"]. The OIG has not issued specific compliance program guidance for distributors. However, the basic program elements discussed in the OIG Pharma Guidance are applicable to distributors as well.

¹⁰⁶ See FSGs 2004 at § 8B.2.1 (The "program shall be reasonably designed, implemented, and enforced so that the program is generally effective in preventing and detecting criminal conduct.").

- Specific individual(s) within the organization shall be delegated day-to-day operational responsibility for the compliance and ethics program. Individual(s) with operational responsibility shall report periodically to high-level personnel and, as appropriate, to the governing authority, or an appropriate subgroup of the governing authority, on the effectiveness of the compliance and ethics program. To carry out such operational responsibility, such individual(s) shall be given adequate resources, appropriate authority, and direct access to the governing authority or an appropriate subgroup of the governing authority.¹⁰⁷

According to the OIG, while there are various ways to demonstrate a company's commitment to compliance, "[e]vidence of that commitment should include the allocation of adequate resources."¹⁰⁸ Therefore, "the compliance measures adopted ... should be tailored to fit the unique environment of the company (including its organizational structure, operations and resources, as well as prior enforcement experience)," and "the compliance officer should have sufficient funding, resources, and staff to perform his or her responsibilities fully."¹⁰⁹ The 2017 compliance program effectiveness guidance documents from both the OIG and DOJ reiterate once more the importance of adequately resourcing the compliance function.¹¹⁰

6.2.1 Attributes

Within the context of a controlled substances compliance program, I would expect a good anti-diversion program for both a manufacturer and a distributor to have the following attributes:

1. **Integration:** The anti-diversion program is integrated into the overall fabric of the organization's corporate compliance program as directed by the Chief Compliance Officer ("CCO"). This can be evidenced by:
 - a. Periodic reports from the Controlled Substances Compliance Team to the CCO, as well as mention of those efforts in the CCO's annual report to the Board of Directors.
 - b. Participation in or periodic input to the Corporate Compliance Committee, either directly or through the appropriate functional leader (e.g., VP of Operations).
 - c. Including an explicit controlled substances compliance expectation within the company's code of conduct.
2. **High-level individual:** The organization assigns responsibility and authority for the anti-diversion program to a relatively high-level individual or group including:

¹⁰⁷ See *id.* at §§ 8B.2.1(a)(2)(A) and(a)(2)(C).

¹⁰⁸ See OIG Pharma Guidance at 23732.

¹⁰⁹ *Id.* at 23732 and 23739.

¹¹⁰ See HCCA-OIG Compliance Effectiveness Roundtable, *Measuring Compliance Program Effectiveness: A Resource Guide*, at 12 (Mar. 27, 2017), available at <https://oig.hhs.gov/compliance/101/files/HCCA-OIG-Resource-Guide.pdf> ["HCCA Effectiveness Guidance"]; U.S. Dep't of Just., Criminal Division, Fraud Section, Evaluation of Corporate Compliance Programs (Feb. 8, 2017), 2-3 <https://www.justice.gov/criminal-fraud/page/file/937501/download> ["DOJ Compliance Evaluation"].

- a. Designating a Vice President level, but no lower than Senior Director, as the highest-ranking person in charge of the anti-diversion program.
 - b. If the program is assigned to someone with additional duties and responsibilities, using the company HR performance review process to assure that he or she understands the success of the anti-diversion program is a key component to their overall compensation package.
 - c. If the program is embedded in an operations group (as opposed to the Office of the Chief Compliance Officer), the creation of an independent reporting line to the CCO.
 - d. Compliance determinations by the high-level individual or group, including customer acceptance/termination and processing of “flagged” orders are appealable only to the CCO or the Compliance Committee, and their decision is final.
 - e. The organization maintains current, accurate organizational charts applicable to the anti-diversion program.
3. **Resources:** The organization provides adequate budget and headcount to carry out the activities of the anti-diversion program effectively.
- a. The budget allows for enough travel funds to conduct onsite visits and investigations and provides some funding to hire outside support as needed.
 - b. If the company leverages indirect reports (e.g., using internal audit staff to conduct customer investigations), using the company HR performance review process to assure that the indirect reports understand that supporting the anti-diversion program is a key component to their overall compensation package.

6.3 Written Standards & Education

Having established written policies and procedures (standards) is fundamental to having an effective compliance program.¹¹¹ Written standards serve as the basis for instructing an organization’s employees not only what tasks need doing (policy) but how they need to accomplish those tasks (procedure). Written standards also are important to ensure consistent outcomes are achieved from the processes the organization utilizes.

While the exact format of policies and procedures vary by organization, there are standard elements common to all policies and procedures, especially in the pharmaceutical industry.¹¹² From a compliance program effectiveness standpoint, scope (do the standards encompass what needs to be addressed?), clarity (are the standards understandable?), accountability (do employees know what they are accountable for and when they must involve the gatekeepers?), and accessibility (can employees find the officially approved standards to read them?) are primary factors in determining whether policies and procedures will be effective.¹¹³

¹¹¹ See FSGs 2004 at § 8B.2.1(b)(1).

¹¹² See, e.g., Margret Amatayakul, *Practical Advice for Effective Policies, Procedures*, 74 J. OF AHIMA.4: 16A-D (Apr. 2003), <http://library.ahima.org/doc?oid=59451#.XBfhIfZFwuW>. For a list of those standard elements, see Appendix B, Figure 1.

¹¹³ See HCCA Effectiveness Guidance, at 3-4; DOJ Compliance Evaluation, at 3-4; see also ISO 9001:2015 (outlining the basic concepts of good document control).

The centerpiece of an organization's collection of written standards, which include company policies and operational procedures, is a core statement of ethical and compliance principles commonly referred to as the "Code of Conduct." The Code of Conduct is the statement of the organization's fundamental principles and values, as well as the expectation that employees will be committed to compliance.¹¹⁴ A Code of Conduct, while not expressly required by the Federal Sentencing Guidelines, nevertheless has been a leading practice since the late 1990's, and is an important mechanism for a company "to communicate effectively its standards and procedures to all employees and other agents" because Codes of Conduct are "publications that explain in a practical manner what is required."¹¹⁵ The OIG also has enshrined the need for a Code in its OIG Program Guidance.¹¹⁶

While established written policies and procedures are critical, just having them is not enough for a compliance program to be effective. An organization's employees must know that those standards exist (communication) and what is expected from each employee (training or education).¹¹⁷ Communicating those standards and expectations is not a "once and done" proposition.¹¹⁸ Because the compliance environment is dynamic with many moving parts (e.g., new hires, new regulations, new policies, new organizational structures, etc.), as well as the fact that people generally need to hear the information more than once to absorb it (e.g., the marketing rule of seven), good compliance functions generally expend significant resources on communication and training.

Furthermore, it is an industry leading practice to require employees to demonstrate mastery of the information being imparted in training (e.g., education) via a test or assessment. Passage of such an assessment provides some modicum of objective evidence that the trainee was effectively trained. The most rigorous programs, especially those using eLearning systems, require the trainee to successfully answer two or three questions after each section to progress and then to pass a final assessment at the end. For "live" or "real-time" training sessions, leading practice is to employ a "sign-in sheet" or some other mechanism to capture attendance. In both cases, the data on attendance and successful completion are normally maintained in an employee's training record, sometimes referred to as their "training jacket," which can be either a digital record or paper file.

6.3.1 Attributes

Within the context of a controlled substances compliance program, I would expect the written standards in a good anti-diversion program for both a manufacturer and a distributor to have the following attributes:

¹¹⁴ See, e.g., OIG Pharma Guidance at 23733.

¹¹⁵ See FSGs 2004 at § 8B.2.1(b)(4).

¹¹⁶ See, e.g., OIG Pharma Guidance at 23733.

¹¹⁷ See FSGs 2004 at § 8B.2.1(b)(4). ("The organization shall take reasonable steps to communicate periodically and in a practical manner its standards and procedures, and other aspects of the compliance and ethics program, to the individuals referred to in subdivision (B) by conducting effective training programs and otherwise disseminating information appropriate to such individuals' respective roles and responsibilities.").

¹¹⁸ *Id.*

1. **Standard Elements:** The written standards incorporate the standard elements common to all policies and procedures in the written standards, including but not limited to Title, Purpose, Scope, Responsibilities, Effective Date, and Revision History.¹¹⁹
 - a. Key terms (e.g., thresholds) are defined either directly in the written standards or via a separate glossary of terms.
 - b. Employees can clearly determine what they will be held accountable for and when they must involve the gatekeepers.
 - c. Where discretion is granted to gatekeepers, the standards define the criteria used in making those decisions.
2. **Document Control:** The written standards are developed, revised and approved utilizing a formal document control process.
 - a. The process, at a minimum, tracks approvals, revisions and the reason for any changes.
 - b. The process ensures that obsolete versions of the standards are withdrawn from use, but maintains withdrawn copies in an archive.
 - c. Depending on the size of the organization, the document control process may be either paper-based or electronic.
 - d. All archived versions of written standards are stored and maintained as essential compliance and business records.
3. **Publication:** The written standards are maintained in a form and location readily accessible to all employees.
 - a. Depending on the size of the organization, publication may be either paper-based (e.g., a manual) or via electronic media such as a company intranet.

Education in a good anti-diversion program would have the following attributes:

1. **Acknowledgment of Standards:** Employees with controlled substances responsibilities acknowledge receipt and having “read and understood” the issued standards in a timely manner (e.g., 10 days).
 - a. Those acknowledgments are collected, tracked, and follow-up occurs for missing or incomplete acknowledgments.
2. **Good Training Practices:** All employee education courses follow good training practices including:
 - a. Depending on the size of the organization, the courses are delivered by face-to-face or eLearning methods.
 - b. Courses follow the principles of good instructional design (e.g., limited duration, information not densely packaged, etc.).¹²⁰
 - c. Session attendance and overall completion are tracked, and follow-up occurs for missing records or incomplete training.

¹¹⁹ For a complete list of standard elements, see Appendix B, Figure 1.

¹²⁰ See, e.g., Presentation Mike Kunkle, *Instructional Design Primer*, (Feb. 6, 2011), <https://www.slideshare.net/MikeKunkle/basic-instructional-design-principles-a-primer> (last accessed Mar. 14, 2019). This presentation is illustrative of the fact that information on good instructional design is widely and readily available.

- d. Successful completion of the course by employees is based on an objective assessment or test demonstrating comprehension of the topics covered.
 - e. Failure to successfully complete a course after 2 or 3 attempts triggers additional follow-up and counseling by the employee's manager and compliance as assisted by HR.
 - f. Every employee has an accurate record of any educational courses completed during their career with the company that is maintained in a readily retrievable format (e.g., a "training jacket" or Learning Management System ("LMS") file).
 - g. The education record, at a minimum, contains course, title, and date, instructor name or LMS file name, and completion outcome (e.g., pass or fail).
 - h. All educational course materials and individual educational records are stored and maintained as essential compliance and business records.
3. **Controlled Substances Education:** Employees with responsibilities for controlled substances compliance complete required education courses.
- a. Education courses for newly hired employees are completed before the new employees can work alone.
 - b. Refresher education courses are conducted on an annual basis.
4. **General Compliance Education:** All employees receive a level of periodic compliance education that includes how to raise questions, as well as reporting issues of suspected misconduct.
5. **Other Educational Methods:** The organization uses other means and methods (e.g., periodic newsletters, "email blasts," etc.) to routinely engage with employees and keep them abreast of impending changes to the anti-diversion program or to solicit employee feedback.

6.4 Monitoring, Auditing & Investigations

Detection or due diligence is at the heart of an effective compliance program.¹²¹ This concept of detection involves three different but interrelated activities (monitoring, auditing, and investigations). Although sometimes used interchangeably, monitoring, auditing and investigations differ in scope and application, but do all ultimately involve looking for anomalous behavior or outliers that need correcting.¹²²

Effective Suspicious Order Monitoring Programs also utilize all three concepts. However, unlike a general corporate compliance program, a SOM program does not simply involve monitoring, auditing and investigating

¹²¹ See FSGs 2004 at § 8B.2.1(a)(1).

¹²² Monitoring is a continuous, real or near-real time activity using established criteria to demonstrate adherence to specific standards. Audits are retrospective, "snapshots in time" to provide assurance that employees are adhering to the required process. Transactional testing is used in audits to verify that the process truly is being followed. Investigations involve examining specific circumstances or individuals to determine if breaches of company policies, procedures or the law have occurred.

internally to ensure the employees are following the prescribed standards, but also applies these activities externally to its customer base as part of the system of controls for preventing diversion.¹²³

By reviewing the DEA regulations and general guidance letters provided to all registrants during the review period, it is possible to get a clear concept of what a successful SOM program should look like. Below is a summarized list of SOM requirements derived from those sources:

1. The customer must be “known” to determine that the customer can lawfully receive the shipment.¹²⁴
2. There must be a designed system.¹²⁵
3. It must be operational.¹²⁶
4. It must identify suspicious orders of controlled substances.¹²⁷
5. Orders can be suspicious because of:¹²⁸
 - a. unusual size;
 - b. substantial deviation from a normal pattern; or
 - c. unusual frequency.
6. Once a suspicious order is discovered,
 - a. the local DEA Field Office must be informed,¹²⁹ and
 - b. the order must be prevented from being filled until it can be ascertained that the order will not be diverted.¹³⁰

Utilized correctly, the establishment of thresholds (a cap on the amount of controlled substances a customer can order in a set period) is an effective way to identify, but not confirm, suspicious orders. Once identified as suspicious, the reasonable, and required steps, include placing a “hold” or “stop notice” on the order to prevent the product from potentially being diverted, immediately notifying the appropriate DEA field office or DEA headquarters or immediately conducting an appropriate investigation to determine if the suspicious order is indeed a potential diversion situation.

Only after the investigation determines that the risk of diversion is not present, can the shipment be processed in the usual course. However, if the investigation determines that there is a risk of diversion, the order must not be filled, and the company should contemplate other appropriate steps for handling future shipment requests. Those steps include refusing to ship any more products to the customer, requiring the customer to provide independent assurance that a diversion situation is not present, or terminating the customer altogether.

¹²³ See 21 C.F.R. §§ 1301.74 (a-b); DOJ Compliance Evaluation at 7 (third party management); James Arnold 6/2013 Presentation at 42-53; Presentation by G. Boggs, *State of Prescription Drug Abuse*, 39 (2013), MCKMDL00336833; see also McKesson, *McKesson Operations Manual for Pharma Distribution, Controlled Substances Monitoring Program*, 56 (Aug. 24, 2011), MCKMDL00000021 (“McKesson’s responsibility is to “Know our Customer.”).

¹²⁴ See 21 C.F.R. § 1301.74(a).

¹²⁵ See 21 C.F.R. § 1301.74(b).

¹²⁶ See 21 C.F.R. § 1301.74(b).

¹²⁷ See 21 C.F.R. § 1301.74(b).

¹²⁸ See 21 C.F.R. § 1301.74(b).

¹²⁹ See 21 C.F.R. § 1301.74(b).

¹³⁰ See DEA 6/12/2012 Letter at 2.

6.4.1 Attributes

A. Distributors

Within the context of a controlled substances compliance program, I would expect the monitoring, auditing, and investigations program for a robust distributor anti-diversion program to have the following attributes:

1. **Know Your Customer:** The distributor has and maintains current granular and specific knowledge about each retail pharmacy customer and their unique circumstances.
 - a. Customer background information:
 - i. Is collected on all customers, including national retail chains, before any product sales are made.
 - ii. Is collected using a standard methodology (e.g., a questionnaire) that balances the need to see patterns and trends amongst similarly situated customers, with the flexibility to capture unique circumstances.
 - iii. Includes more than DEA and state Board of Pharmacy licenses to include available internet or commercially obtainable information (e.g., GOOGLE searches, Dun & Bradstreet reports, IMS data, etc.).
 - iv. Includes dispensing data and largest prescribers.
 - v. Includes whether the relationship is primary (e.g., exclusive) or secondary.
 - vi. Includes the customer's prior overall compliance history that is not limited to just controlled substances.
 - vii. Uses a risk-adjusted process for periodically re-evaluating customers considering changed circumstances. Those risk factors include the indicators of diversion provided by the DEA as well as changes in control (e.g., merger, acquisition, the sale of a business) or customer profile (e.g., new pain clinic in the territory served).
 - viii. Obtains references from the primary distributor when known.
 - ix. Is kept current and updated on a regular, periodic basis.
 - b. Customer background information is evaluated for completeness and accuracy.
 - i. Submissions of inaccurate or incomplete information are grounds for immediate disqualification or termination.
 - ii. Refusal to provide requested background information is grounds for immediate disqualification or termination.
 - iii. In the case of the large retail pharmacy chains (e.g., CVS, Walgreens, and Rite Aid) defines "customer" in terms of the individual retail pharmacy location and not just the national chain.
 - c. Customers are evaluated and approved or denied based upon submitted background information and any other due diligence conducted by the organization.
 - i. Evaluations occur under established criteria, which, at a minimum, incorporate any guidance from the DEA.
 - ii. "Red flags" such as being a secondary supplier or customer being recently terminated by another distributor trigger a thorough investigation including a site visit by a trained investigator.
 - iii. Outcomes are clear and well-documented.

- d. Customer site visits are routinely and periodically performed even after an initial site visit.
 - i. Site visits are performed by individuals trained in diversionary behaviors.
 - ii. If sales personnel are utilized to perform site visits, steps are taken to minimize conflicts of interests (e.g., using out-of-territory personnel).
 - e. Customer files are stored and maintained as essential compliance and business records.
2. **Thresholds:** The organization uses threshold calculations based on dosage units to identify “suspicious orders.”
- a. Thresholds are customer-specific and set using the background information obtained and maintained by the organization in accordance with the organization’s written standards.
 - b. Like customers are grouped together with as much granularity as possible (e.g., business activity, purchasing patterns, total prescriptions, geographic location, size of territory served).
 - c. Thresholds are calculated based on multiple criteria using a documented, validated statistical formula that considers, at a minimum, the following items:
 - i. Customer group;
 - ii. Order size, patterns, and frequency, both individually and of the group; including orders being filled by other distributors;
 - iii. Dispensing data, both individually and of the group;¹³¹
 - iv. Geographic territory and population served;
 - v. Product formulation (active ingredient and dosage) as well as the diversion potential; and
 - vi. Legitimate, medically necessary, dosage unit levels developed based upon the approved indications for use and without regard to current opioid purchasing patterns.
 - d. A minimum of 12-months of relevant and complete historical data is used without “cherry picking” the most favorable data.
 - e. Actual thresholds and the threshold calculation methodology are not shared with customers.
 - f. Thresholds are binding until an approved threshold exception or adjustment is granted.
3. **Threshold Exceptions or Adjustments:** Any threshold exceptions or adjustments rarely are made when viewed by the individual customer or the group, as well as a simple review of request frequency.
- a. Threshold exceptions or adjustments are made by a committee of individuals with anti-diversion experience and training in accordance with the organization’s written standards.
 - b. All threshold exceptions or adjustments are supported by verifiable objective evidence that is documented in writing.
 - c. Threshold exceptions and adjustments are tracked and trended on both a short-term (e.g., weekly or monthly) and long-term (e.g., quarterly or annual) basis.
 - d. All threshold exceptions or adjustments, including any records of approvals or denials, are stored and maintained as essential compliance and business records.

4. **Taking Action:**

¹³¹ With the appropriate safeguards to protect patient identifiable health information under HIPPA and other relevant privacy standards.

- a. All orders which exceed the customer-specific threshold are deemed “suspicious” and reported to the DEA within one week unless it is determined that there are no reasons to suspect that a customer is engaging in diversion; for example, a clerical mistake (e.g., “fat-finger” orders).
 - b. All orders exceeding thresholds are held and not shipped. No order “cutting” is permitted.
 - c. No orders exceeding thresholds may ship until a thorough independent investigation is completed based on verifiable objective evidence demonstrates that diversion is unlikely to occur, and the findings are reviewed and approved.
 - i. The use of the Corporate Headquarter staff of the large retail pharmacy chains (e.g., CVS, Walgreens, and Rite Aid) as the “lead investigator” is not permitted to maintain independence.
 - d. No future orders involving the same active ingredient are processed or shipped until the thorough independent investigation is completed based on verifiable objective evidence that demonstrates diversion is unlikely to occur and the findings are reviewed and approved.
5. **Audits:** The distributor’s internal audit team, or an appropriately qualified third-party (e.g., the company’s external auditors) conducts periodic, regular audits of the distributor’s anti-diversion program including transactional testing of a statistically relevant sample of retail pharmacy customers.
- a. All customer supply contracts contain the appropriate “right-to-audit” clause.
 - b. All audits are conducted in accordance with written standards.
 - c. Audits are conducted on a risk-adjusted basis.
 - d. Audit findings and corresponding management responses are tracked and trended.
 - e. Repeat audit findings are escalated to the organization’s Chief Executive Officer and the Audit Committee of the Board of Directors.

B. Manufacturers

For a manufacturer’s anti-diversion program, I would expect to see:

1. **Know Your Customer:** The manufacturer has and maintains current granular and specific knowledge about each distributor of its controlled substances and their unique circumstances including all the information outlined in the distributor section above.
 - a. Distributor site visits are undertaken to review the distributor’s anti-diversion controls both at initiation of the relationship and then periodically on a risk adjusted basis thereafter (see Audits section below).
 - b. Utilize, where appropriate, information derived from chargeback data.
2. **Individual Retail Pharmacy Activity:** Like the distributor thresholds outlined above, the manufacturer establishes ordering levels for specific pharmacies, which if exceeded trigger the manufacturer to be concerned that the orders are “suspicious,” and that action is needed.
 - a. Where appropriate, information obtained through the manufacturer’s sample accountability (e.g., PDMA) program is factored into the controlled substances monitoring program.
 - b. Wherever possible, the manufacturer leverages synergies (people, process and technology) between the sample accountability and controlled substances compliance programs.

3. **Taking Action:** When the manufacturer gains knowledge of retail pharmacies placing “suspicious orders” or otherwise engaging in diversionary behavior (e.g., serving questionable prescribers), the manufacturer takes the following actions:
 - a. The manufacturer notifies and provides details of the suspicious activity to both the DEA and the distributor.
 - b. The manufacturer demands the distributor, and any secondary distributor if known, follow-up and take appropriate action regarding the highlighted pharmacies.
 - c. The manufacturer maintains contact with the distributor, and any secondary distributor if known, requiring them to provide details on the outcome of any investigations including actions taken by the distributor(s) against the pharmacies.
4. **Audits:** The manufacturer conducts both routine and “for cause” audits of those distributors’ anti-diversion programs.
 - a. All customer supply contracts contain the appropriate “right-to-audit” clause.
 - b. Routine audits are conducted on a risk-adjusted basis.
 - c. All audits are conducted in accordance with written standards.
 - d. Audit findings and corresponding management responses are tracked and trended.
 - e. Repeat audit findings are escalated to the organization’s Chief Executive Officer and the Audit Committee of the Board of Directors.

6.5 Corrective Actions & Risk Assessments

Once non-compliant conduct, whether criminal or not, has been detected by monitoring and confirmed to have occurred through investigation, the organization is expected to determine and implement changes to avoid either a continuation of the underlying conduct, or to prevent a new occurrence from arising.¹³² As the OIG elaborated:

Violation of a pharmaceutical manufacturer’s compliance program, failure to comply with applicable federal or state law, and other types of misconduct threaten the company’s status as a reliable, honest, and trustworthy participant in the health care industry. **Detected but uncorrected misconduct can endanger the reputation and legal status of the company.**

Consequently, upon receipt of reasonable indications of suspected noncompliance, it is important that the compliance officer or other management officials immediately investigate the allegations to determine whether a material violation of applicable law or the requirements of the compliance program has occurred and if so, take **decisive steps** to correct the problem.¹³³

¹³² See FSGs 2004 at § 8B.2.1(b)(7) (“After criminal conduct has been detected, the organization shall take reasonable steps to respond appropriately to the criminal conduct and to prevent further similar criminal conduct, including making any necessary modifications to the organization’s compliance and ethics program.”).

¹³³ See OIG Pharma Guidance at 23742 (emphasis added); HCCA Effectiveness Guidance at 50-51 §§ 7.43-7.54.

The DEA regulations also embody the corrective action concept.¹³⁴ Identified corrective actions need to be documented and monitored to ensure they are implemented. This is especially true for complex corrective actions that often span many weeks or months to accomplish.

For a compliance program to be effective, just correcting errors, omissions and breaches are not enough. Organizations also need a documented process to conduct risk assessments.¹³⁵ The risk assessment process captures changes as various risks morph (e.g., the emergence of internet pharmacies dispensing controlled substances), as well as what the organization is doing to address or mitigate those risks and to assess whether those activities are working.¹³⁶

Typically, when companies first embark on establishing a compliance program, they engage in a risk assessment, more often referred to as a “gap analysis.” This gap analysis provides the compliance program designers with crucial information on what needs to be addressed.

However, risk assessments normally are not a single event. The risk assessment process envisioned by the FSGs is a true, repeatable process that should occur at regularly scheduled intervals. Therefore, while the gap analysis is usually done at the beginning, the formal risk assessment process frequently is established after the basic compliance program framework is in place.

6.5.1 Attributes

Within the context of a controlled substances compliance program, I would expect the corrective action and risk assessment processes for both a robust distributor and manufacturer anti-diversion program to have the following attributes.

1. **Corrective Actions:** The organization has a formal, documented corrective action process, and applies that process to the anti-diversion program:
 - a. All program deficiencies are documented regardless of source (e.g., internal or external audits, internal or external assessments, regulatory inspections, regulatory guidance, and industry standards).
 - b. For every documented deficiency, a plan for correction, which details the remedy, employees’ responsible for making the corrections and plan milestones, is developed.
 - c. The final approved plans are collected and tracked with corresponding updates to the Compliance Committee, Senior Management and if warranted the Audit Committee of the Board of Directors.
 - i Whenever possible, corrective action documentation and tracking are incorporated into the organization’s electronic Governance, Risk and Compliance or e GRC system (e.g., Archer).

¹³⁴ See, e.g., 21 C.F.R. § 1301.71(c).

¹³⁵ See FSGs 2004 at § 8B.2.1(c) (“In implementing subsection (b), the organization shall periodically assess the risk of criminal conduct and shall take appropriate steps to design, implement, or modify each requirement set forth in subsection (b) to reduce the risk of criminal conduct identified through this process.”).

¹³⁶ See DOJ Compliance Evaluation at 4-5, Topic 5; HCCA Effectiveness Guidance, at 15 §§ 2.56-2.62.

- d. Plan timelines have a finite time limit (e.g., no more than 12 months)
 - i If additional time is necessary to complete the corrective action, a new plan is submitted and approved.
 - ii Senior Management and Compliance approval is required for any extensions.
- e. Individual accountability is managed through the standard HR performance appraisal process.
- f. Corrective action items are only closed upon independent verification that the planned corrections are complete and functioning as intended.

2. **Risk Assessments:** The organization maintains a formal, documented risk assessment process to evaluate legal and compliance risks to the organization's anti-diversion program. A robust process includes, but is not limited to, the following elements:

- a. The process evaluates both internal and external risks to the anti-diversion program.
- b. The process leverages data from all available sources, including but not limited to:
 - i Budgets;
 - ii Headcount;
 - iii Exit interviews;
 - iv Employee surveys;
 - v Investigation results;
 - vi Audit results;
 - vii Corrective actions;
 - viii Commercial benchmarking data (e.g., IMS data);
 - ix Regulatory inspections; and
 - x Enforcement actions.
- c. A risk assessment review occurs at defined intervals, but no less than annually.
- d. The risk assessment output is documented and is:
 - i Disseminated widely to management, compliance, legal, internal audit and those responsible for the anti-diversion program.
 - ii Maintained in a readily digestible format such as a "heat map."
 - iii Incorporated, whenever possible into the organization's eGRC system.
 - iv Used to develop further corrective actions, audit planning, budget and headcount increases, customer monitoring efforts, etc.
- e. Previous risk assessment outputs are maintained and utilized for benchmarking and trending purposes to show improvement or decline in the effectiveness of the anti-diversion program.

6.6 Accountability - Consistent Enforcement

Accountability also is a fundamental element of an effective compliance program. Under the FSGs, there are two intertwined provisions that apply in this context. The first involves consistent enforcement of compliance

and ethical standards, including government requirements, otherwise known as discipline.¹³⁷ The second involves being careful with the delegation of substantial authority, otherwise known as “avoiding bad actors.”¹³⁸

6.6.1 Discipline

In the case of consistent enforcement, the FSGs succinctly notes “[a]dequate discipline of individuals responsible for an offense is a necessary component of enforcement.”¹³⁹ The OIG Compliance Program Guidance goes further stating:

Intentional and material noncompliance should subject transgressors to significant sanctions. Such sanctions could range from oral warnings to suspension, termination or other sanctions, as appropriate. **Disciplinary action also may be appropriate where a responsible employee’s failure to detect a violation is attributable to his or her negligence or reckless conduct.**¹⁴⁰

6.6.2 Avoiding “Bad Actors” – Employees or Customers

The FSGs also requires organizations to “use reasonable efforts” to ensure that “with the substantial authority personnel of the organization any individual whom the organization knew or should have known through the exercise of due diligence, has engaged in illegal activities or **other conduct inconsistent with an effective compliance and ethics program**” are not placed in a position to cause harm through their non-compliant actions.¹⁴¹

The FSGs defines “substantial authority personnel” as:

individuals who within the scope of their authority exercise a substantial measure of discretion in acting on behalf of an organization. The term includes high-level personnel of the organization, individuals who exercise substantial supervisory authority (e.g., **a plant manager, a sales manager**), and any other individuals who, although not a part of an organization’s management, nevertheless exercise substantial discretion when acting within the scope of their authority (e.g., an individual with **authority** in an organization **to negotiate or set price levels** or an individual authorized to negotiate or approve significant contracts).¹⁴²

While the FSGs focuses primarily on organizational and employee accountability, the CSA and its implementing regulations focus on customer behaviors. Embedded within the concept of identifying suspicious orders and having effective diversion controls is the common sense proposition that if an order is initially

¹³⁷ See FSGs 2004 at § 8B2.1(b)(6).

¹³⁸ See FSGs 2004 at § 8B2.1(b)(3).

¹³⁹ See FSGs 2004 at § 8B2.1, Application Note 5.

¹⁴⁰ See OIG Pharma Guidance at 23742 (emphasis added).

¹⁴¹ See FSGs 2004 at § 8B2.1(b)(3) (emphasis added).

¹⁴² See FSGs 2004 at § 8A1.2, Application Note 3(c) (emphasis added).

flagged as suspicious using the criteria in the DEA regulations (unusual size, pattern, frequency), the distributor must not ship that order or any similar controlled substances order to that customer until the distributor determines whether or not there is likelihood the shipment is being diverted.¹⁴³ To do otherwise, potentially allows a diversionary situation to continue, which is the opposite of preventing diversion.¹⁴⁴ In other words, the distributor is expected to impose discipline on its customers when the distributor becomes aware of customers that are placing suspicious orders.

6.6.3 Attributes

1. **Employees:** The organization (both distributors and manufacturers) maintains a robust screening (background check) process and a disciplinary system that includes appropriate sanctions up to and including termination.
 - a. Employees alleged to have violated any anti-diversion requirements are immediately removed from any further responsibilities involving controlled substances until cleared by a thorough independent investigation demonstrating that no violation occurred based on verifiable objective evidence.
 - b. Employees who violate the requirements of the organization's anti-diversion program are subject to appropriate disciplinary sanctions.
 - c. Disciplinary sanctions are routinely and consistently enforced regardless of an employee's level in the organization or previous job performance.
2. **Distributor Customers:** Retail pharmacy customers, failing to comply with any requirements of the distributor's anti-diversion program (e.g., providing incomplete or inaccurate information) are subject to immediate disqualification or termination.
 - a. This requirement is explicitly stated in all customer supply contracts.
 - i. Contracts contain a "for cause" immediate termination provision, which includes being non-compliant.
 - b. Disqualifications and terminations are routinely and consistently enforced regardless of a customer's prior purchasing history.
 - c. Any pending shipments are immediately canceled.
 - d. Disqualified or terminated customers are not eligible for reinstatement until a thorough audit is conducted and any corrective actions by the customer are verified via objective evidence demonstrating that the customer has effectively corrected all issues underpinning the disqualification or termination.

¹⁴³ DEA 2/7/2007, 12/27/2007, 6/12/2012 Letters at 2; *see also Southwood Pharmaceuticals, Inc.*, Revocation of Registration, 72 Fed. Reg. 36487, 36500 (Jul. 3, 2007) (Holding the distributor accountable for not stopping shipments to customers it should have known were placing suspicious orders including those customers DEA told the distributor were engaging in suspicious ordering).

¹⁴⁴ *See* 72 Fed. Reg. at 36500 ("In short, the direct and foreseeable consequence of the manner in which Respondent conducted its due diligence program was the likely diversion of millions of dosage units of hydrocodone. Indeed, it is especially appalling that notwithstanding the information the Respondent received from both this agency [DEA] and the pharmacies, it did not immediately stop distributing hydrocodone to any of the pharmacies. Moreover, in several cases, Respondent actually distributed even larger quantities of the drug to them.")

- i. Reinstatement of disqualified or terminated customers is reviewed and approved by either the CCO or Compliance Committee.
 - e. Notices of customer disqualifications or terminations are communicated as soon as possible to the distributor's sales representatives.
 - i. The distributor adjusts sales representative compensation plans to remove any negative impact from disqualification or termination.
3. **Manufacturer Customers:** Distributor customers of the manufacturer, which distribute the manufacturer's prescription opioid products are subject to appropriate disciplinary sanctions up to and including termination of the relationship.
- a. This requirement is explicitly stated in all customer supply contracts.
 - i. Contracts contain a "for cause" immediate termination provision, which includes being non-compliant either with the manufacturer's anti-diversion requirements or when cited by the DEA.
 - b. Contracts allow for the immediate cessation of chargebacks for prescription opioid products to non-compliant retail pharmacy customers.

6.7 Manufacturer – Prescriber Relationship

Opioid manufacturers within the DEA's "closed-loop" system, unlike distributors, also are uniquely positioned to observe prescriber behaviors. This occurs because the manufacturers' field forces make routine sales calls on prescribers' offices. Thus, the field forces can be exposed to some of "red flag" indicators such as overly full waiting rooms, young patients, people nodding off in the waiting room, etc.¹⁴⁵ Put another way, things that "if you were to walk into a doctor's office would give you pause and would make you turn around and walk out."¹⁴⁶ The same is true for information obtained from other sources such as IMS data, or media reports.

Given this unique vantage point, the prudent and responsible manufacturer should instruct and require its sales representatives, and in-house field support and marketing personnel, to provide any observations of potential diversionary behavior to their in-house Compliance Department for further evaluation and potential action. As Acting Administrator Rosenberg noted in the Masters Pharmaceutical proceedings, "a registrant cannot ignore information it obtains that raises a suspicion not only with respect to a specific order, but also as to the legitimacy of a customer's business practices" or more specifically, "a registrant cannot claim that it ... has an effective suspicious orders monitoring program when it ignores information it has acquired which raises a substantial question as to the legitimacy of a customer's dispensing practices."¹⁴⁷ While the company needs to act with care to be objective (which is true for every compliance investigation), "turning a blind eye" is not an option.

¹⁴⁵ See Scott Glover and Lisa Giron, OxyContin maker closely guards its list of suspect doctors, LOS ANGELES TIMES (Aug. 11, 2013), <https://www.latimes.com/local/la-xpm-2013-aug-11-la-me-rx-purdue-20130811-story.html>.

¹⁴⁶ See *id.* (quoting Robin Abrams, attorney for Purdue Pharma and a former federal prosecutor specializing in federal healthcare fraud).

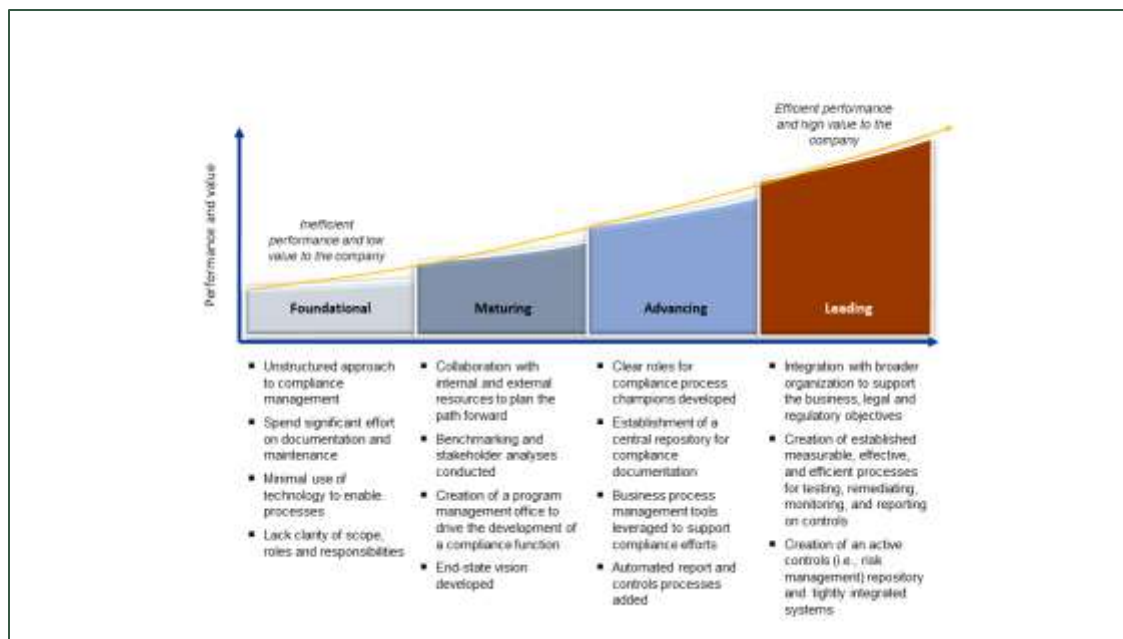
¹⁴⁷ See 80 Fed. Reg. 55418, 55478 (Sept. 15, 2015).

Upon receipt of this information, the Compliance Department, or other experienced investigators, should conduct an appropriate investigation to determine the validity of the information, using all available sources of information (e.g., the internet, IMS data, etc.), and if confirmed, formulate an appropriate action plan. Depending on the weight of the evidence gathered, that plan can range from conducting further comprehensive monitoring activities to refusing to make further sales calls on the suspect practices to, in the most egregious cases, providing the information to the appropriate authorities, including the DEA and State Medical Boards.

7 Measuring What Good Looks Like

After defining “what good looks like” the next step is to measure it. Measuring compliance effectiveness or “what good looks like” is not simply a matter of taking the attributes and applying a statistical, or even a generally recognized standard scoring methodology, as one does not exist. The best approximation of a standardized scoring model is the compliance maturity and program effectiveness model, which outlines the typical evolutionary pathway most compliance organizations follow.

Figure 2– Compliance Maturity & Program Effectiveness Model



The model sets out a framework outlining what characteristics distinguish a compliance function that is just starting out or where management does not embrace the value of the program from one which is fully embedded into company operations and where management clearly recognizes the value that strong compliance provides. Since the levels of maturity directly correlate to the effectiveness of the compliance program, this model also provides a way to level-set among companies in the same field (e.g., pharmaceutical distribution). Overall, most companies focus on and strive to be in either the advancing or leading categories.

This report first analyzes each distributor’s and a single manufacturer’s overall compliance efforts surrounding controlled substances by starting with suspicious order monitoring. For each company, the analysis focuses on answering two questions. The first question is whether objective evidence exists supporting that the company

being reviewed worked to establish a suspicious order monitoring system, as well as controlled substances and corporate compliance programs. Only if there is evidence that the company did so is the second question relevant.

The second question is whether there is objective evidence showing that the company met its three-prong program effectiveness requirement by (a) having a program that prevents and detects criminal conduct by an organization's employees and (b) maintaining effective controls against diversion, including (c) maintaining and operating an effective system to identify, hold, investigate and report suspicious orders of controlled substances.

PART IV: Report Overview



8 Executive Summary

8.1 Group 1 Distributors

The Group 1 (“G1”) distributors (also known as the “Big Three”), on a national basis, account for 85% of the national drug supply.¹⁴⁸ Although each G1 distributor’s detailed approach to both corporate compliance and anti-diversion controls for controlled substances was reviewed separately, there are common threads that unite all three companies.

¹⁴⁸ See W.Va. Red Flags Report at 7.

First, even though the applicable standards for controlled substances and corporate compliance were established in the early 1970s and 1990s respectively, all three distributors did not establish their own programs until years later. Nor did they make their controlled substances efforts part of their overall corporate compliance programs in a meaningful way.

Second, prior to having significant pressure brought to bear on them by the DEA for non-compliance, none of the three companies made more than token efforts to implement a compliance program to detect and prevent the shipment of prescription opioid products into an illicit market, let alone to fulfill the required legal and regulatory obligations. In so doing, all three failed to manifest good corporate responsibility by not undertaking the reasoned, prudent and careful measures expected of those handling prescription opioid products.

Third, despite entering into settlement agreements to operate effective anti-diversion and SOM programs and claiming to be working with the DEA, each distributor designed, implemented and operated their controlled substances program in such a manner as to avoid classifying customer orders as “suspicious” in order to avoid having to stop suspect opioid shipments to customers. Even after the DEA directed them to act, every G1 distributor failed to make the necessary changes in order to achieve a robust and effective compliance function in accordance with the values, principles and societal expectations for those involved in distributing prescription opioids.

Fourth, when confronted with objective evidence of inappropriate opioid ordering, each company still did not make reasonable inquiries as to “why” it was occurring. By doing so, all three companies studiously avoided having to address customer behaviors they knew or should have known were inappropriate and likely diversionary.

In short, throughout the review period, the big three distributors failed to act responsibly to undertake the reasoned, prudent and careful measures expected of those handling prescription opioid products even while acknowledging the exponential increase in opioid usage. Customer relationships simply trumped compliance. As a result, on the compliance maturity and program effectiveness model, all three score no higher than the midpoint of the foundational level, which is unacceptable and unreasonable given how long these standards have existed, the resources available to each company and the evidence that this class of pharmaceutical product (e.g., opioids) have a high risk of being diverted and a great propensity to cause harm when used improperly.

8.2 Group 2 Distributors

The Group 2 (“G2”) distributors are large, national pharmacy or retail chains, which have embedded distribution operations that supply only their own pharmacies. Although pharmacies are part of the overall controlled substances “closed loop” system,¹⁴⁹ this review is focused on controlled substances compliance only in the context of those internal distribution operations.

While there is no uniformity among the G2 group regarding whether they distributed both Schedule II or Schedule III opioid medicinal products or just Schedule III products,¹⁵⁰ all group members ultimately

¹⁴⁹ See Discussion *infra*.

¹⁵⁰ For example, Walgreens internally distributed both Schedule II and III opioids, while CVS only handled Schedule III opioids internally.

outsourced the distribution of opioid products to their retail locations upon the reclassification of hydrocodone combination products (“HCPs”) from Schedule III to Schedule II in October 2014.¹⁵¹ After that date, opioid products were provided to G2 retail pharmacies via Group 1 (“G1”) distributors sometimes augmented by other independent distributors serving as secondary suppliers.

Although the G2 distributors were like the G1 distributors in that profits trumped compliance, the G2 companies focused most of their anti-diversion efforts on protecting their retail pharmacy business. Consequently, the need for distribution centers to maintain reasoned, prudent and careful measures to prevent opioid diversion (e.g., a SOM program) which is expected of those handling prescription opioid medicines was treated as an afterthought, if it was recognized at all. Furthermore, distribution center anti-diversion efforts tended to focus on losses and thefts (e.g., loss prevention), rather than on whether they were shipping suspicious orders.

Again, even though applicable standards for corporate compliance and controlled substances anti-diversion programs were established in the early 1970s and 1990s respectively, it was not until the 2008-2009 timeframe that they undertook any meaningful efforts to meet their legal, regulatory and societal obligations. Nor did they make their controlled substances efforts part of their overall corporate compliance programs in a meaningful way.

The G2, for the most part, made only token efforts to implement a compliance program to detect and prevent the shipment of prescription opioid products into an illicit market and then only when the DEA directed them to. Thus, both companies failed to manifest good corporate responsibility by not undertaking the reasoned, prudent and careful measures expected of those handling prescription opioid products. For example, although all the G2 distributors had ready access to their own dispensing data, none of them tried to incorporate that information into their anti-diversion programs.

In addition, since retail pharmacies represent a significant profit center for the G2 group, those internally who were charged with controlled substances compliance invested substantial time and resources trying not to classify excessive pharmacy orders as “suspicious,” so as not to disrupt product supply. This constituted an inherent conflict of interest that elevated profits over compliance. In short, throughout the review period, the G2 distributors failed to act responsibly to undertake the reasoned, prudent and careful measures expected of those handling prescription opioid products even while acknowledging that there was an exponential increase in opioid usage.

On the compliance maturity and program effectiveness scale, the G2 companies are barely starting into the foundational level, and while the model does not have a remedial level, if it did, that is where they would be found. Their behavior is unreasonable given the fact that they understood that (a) opioid products have a high risk of being diverted and a great propensity to cause harm when used improperly and (b) they simultaneously occupied two positions in the “closed loop” system (e.g., dispenser and distributor).

¹⁵¹ See 79 Fed. Reg. 49661 (Aug. 22, 2014).

8.3 Manufacturer Group

Although only one manufacturer was examined in this review (Mallinckrodt) given its market presence, I believe that it is possible to compare the anti-diversion efforts of manufacturers and distributors. From 1996 to 2017, Mallinckrodt was a leading manufacturer of generic opioid products, selling over \$18 billion in opioid products.¹⁵² For Summit and Cuyahoga Counties, Mallinckrodt via one of its subsidiaries was the largest supplier of opioid products. Thus, between 2006-2014, Mallinckrodt shipped more opioid products – 26.7% of the total or 2,363,328,618 MMEs – than any other manufacturer.¹⁵³

For manufacturers, while the anti-diversion and corporate compliance standards are the same, as well as many of the controls (e.g., thresholds, due diligence, qualifying new customers, etc.), a manufacturer's anti-diversion program is geared towards looking at the distributor's customers, for which it has access to a large amount of information, in order to assure itself that the distributors are taking the necessary steps to avoid providing opioid products to pharmacies engaging in diversionary behavior.

In the same vein, the issues seen with Mallinckrodt's anti-diversion efforts mirror those seen with both the G1 and G2 distributors. First, even though the applicable standards for controlled substances and corporate compliance were established in the early 1970s and 1990s respectively, Mallinckrodt did not apply serious efforts to establish a credible anti-diversion program until 2008.

Second, as was seen with both the G1 and G2 distributors, Mallinckrodt was determined to only do the bare minimum as expressed by the DEA, which resulted in token efforts to implement a compliance program to detect and prevent the shipment of prescription opioid products into an illicit market, let alone to fulfill the required legal and regulatory obligations. Mallinckrodt, in so doing, failed to manifest good corporate responsibility by not undertaking the reasoned, prudent and careful measures expected of those manufacturing and selling opioid products.

Third, like all the distributors examined, Mallinckrodt designed, implemented and operated its SOM program in such a manner as to avoid classifying customer orders as "suspicious" in order to avoid having to stop suspect opioid shipments to customers. Even after the DEA provided Mallinckrodt with explicit directions such as "If you think it is suspicious, don't fill it,"¹⁵⁴ Mallinckrodt selectively interpreted those explicit directives in such a manner so as not to impact its continued distribution of opioid products, although doing so rendered its anti-diversion program ineffective. Thus, Mallinckrodt repeatedly failed to make the necessary changes to its anti-diversion practices in order to achieve a robust and effective compliance function in accordance with the values, principles and societal expectations for those involved in distributing prescription opioids. Also, when confronted with objective evidence of inappropriate opioid ordering, Mallinckrodt still did not make reasonable inquiries as to "why" this ordering was occurring. By doing so, Mallinckrodt studiously avoided having to address customer behaviors it knew or should have known were inappropriate and likely diversionary.

¹⁵² See January 30, 2019 Mallinckrodt Response to Interrogatory No. 33 & Ex. E.

¹⁵³ See Supplemental Expert Report of Craig J. McCann, Ph.D., CFA, ¶ 13 & Table 1 (April 15, 2019).

¹⁵⁴ See Email from B. Ratliff to J. Rausch, *et al.*, Suspicious Order Monitoring, (Apr. 1, 2008) (Karen Harper was copied on this email), MNK-T1_0000268860.

Finally, comments by company employees such as opioids are ‘just like Doritos; keep eating, we’ll make more’”¹⁵⁵ exemplify that although Mallinckrodt espoused company values of being patient-centric and operating with integrity, these values were platitudes. The previous comments also show that Mallinckrodt apparently was indifferent to any negative societal impact flowing from its actions. Consequently, while Mallinckrodt publicly stated that this behavior “is antithetical to everything that Mallinckrodt stands for and has done to combat opioid abuse and misuse,”¹⁵⁶ Mallinckrodt’s current Chief Commercial Officer and previous President of the company’s generics division, also repeatedly refused under oath to express any opinions about the behavior, let alone condemn it as being “antithetical” to Mallinckrodt’s corporate values.¹⁵⁷

As a result of all the issues highlighted above, on the compliance maturity and program effectiveness scale, Mallinckrodt’s program falls at the extreme lower end of the foundational level.

8.4 An Integrated Ecosystem

The DEA “closed loop” system for the distribution of opioid products is no different from a biological ecosystem in which each group (“species”) has a distinct role to play, and if a group or even some members of that group fails to perform their allotted role, the foundation of the ecosystem is threatened.

As previously discussed, distributors and manufacturers both play a role in the controlled substances ecosystem. For the review period, based upon the individual company information I reviewed, the companies reviewed here did not maintain and operate the type of anti-diversion programs that are expected of prudent and socially responsible companies handling this type of high-risk product.

Furthermore, manufacturers and distributors also are corporate-entities focused on shareholder return. As such, there is limited incentive for each to use the information which it has access to, either to enhance its own controlled substances compliance program or to share information about improper pharmacy behavior with others in the opioid supply chain. Thus, a predictable outcome of the failure of manufacturers and distributors to maintain an effective anti-diversion program is that it allows opioid product dispensers (i.e., retail pharmacies) to use the weaknesses in each company’s program to their advantage to maintain a consistent, dependable supply of opioid products to meet physician and patient (i.e., customer) demand whether legitimate or not.

Therefore, because the “closed loop system” is an ecosystem, any examination should look at the operation of the full ecosystem as well as the individual parts. Euclid Family Pharmacy and CVS stores 3322 and 4800 provide excellent examples to do so.

¹⁵⁵ See Associated Press, *Suit: US Drug Agency Deemed Firm ‘Kingpin’ in ‘Drug Cartel,’* THE NEW YORK TIMES (Apr. 1, 2019), <https://www.nytimes.com/aponline/2019/04/01/us/ap-us-opioid-lawsuit-tennessee.html>.

¹⁵⁶ *Id.*

¹⁵⁷ See Hugh O’Neill Deposition, 152:15-153:4; 155:23-158:15; 166:10-169:15; 171:10-175:19 (Mar. 13, 2019).

A. Euclid Family Pharmacy

Located in Cuyahoga County Ohio, the Euclid Family Pharmacy sits in the lobby of a medical building that serves a suburb of Cleveland with a population of approximately 50,000 people.¹⁵⁸ Since 1995, Euclid has been owned and operated by Timothy Williams, a licensed Ohio pharmacist.

As a result of being in the medical building, a significant portion of Euclid's business comes from the pain clinics co-located in the building.¹⁵⁹ As a result of Euclid's advantageous location, pharmaceutical manufacturers have long targeted Euclid for its opioid product sales potential.

For example, in 2006, Endo Pharmaceuticals listed Euclid as one of the "high opioid potential" pharmacies in the region.¹⁶⁰ That list was circulated to Endo's sales representatives with specific instructions about how they could leverage key information about physicians, pharmacies, and distributors to maximize opioid product sales. Endo specifically instructed its representatives to ask physicians "to assist your stocking effort by sending a note or making a call to the pharmacy that they send their patients to fill their opioid prescriptions." Endo also instructed the representatives to follow up with pharmacies "to ensure that the drug is ordered." and to "inquire as to who is the primary wholesaler and back-up wholesaler the pharmacy uses and record that information for your information at a later time."¹⁶¹

Purdue Pharma L.P., engaged in a similar effort in 2008 as it launched a new promotional program for OxyContin.¹⁶² The Purdue program allowed customers to redeem OxyContin Savings Cards, which had the effect of increasing the period of time a patient would use the drug.¹⁶³ Euclid Pharmacy was one of the pharmacies which participated in the OxyContin Savings Card program, and by 2009, the effects were evident as Purdue's sales of oxycodone and hydrocodone notably increased.¹⁶⁴ Although the sales increases are notable on their own, the increases are more troubling when weighed against the fact that in 2006, Mr. Williams told a reporter for that he believed OxyContin was more addictive than other prescription opiates.¹⁶⁵

[REDACTED]

¹⁵⁸ [REDACTED]

¹⁵⁹ *Id.*

¹⁶⁰ Endocet is the brand name for the combination of oxycodone and acetaminophen.

¹⁶¹ See Email from Chris Speaker to Teresa Leigh, Pharmacy Calls, (Aug. 25, 2006) (enclosing an email from Mike Weber to All Pharma Representatives) ENDO-OPIOID_MDL-02776076.

¹⁶² OxyContin is a controlled release version of oxycodone that Purdue manufactured. OxyContin is the brand name.

¹⁶³ See <https://www.wbur.org/commonhealth/2019/02/01/purdue-oxycontin-savings-cards> (last visited March 25, 2019).

¹⁶⁴ See Spreadsheet, March 2009 OxyContin Savings Cards Redemption, PPLPC004000194244; ABDCMDL00170149.

¹⁶⁵ See Purdue News Clips Report, (undated), PPLPC039000161600 at PPLPC039000161640.

¹⁶⁶ [REDACTED]

Euclid also was associated with a local healthcare and pain management clinic and most of its patients were coming from a doctor's office or clinic in the building.¹⁶⁸

██████ Hence, the original customer questionnaire was incomplete, as it does not indicate the required Director of Regulatory Affairs review occurred.

[REDACTED]

¹⁶⁷ See Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated Mar 25, 2019 at 318 (Oxycodone Distribution to Euclid Family Pharmacy).

¹⁶⁹ *Id.*

¹⁷¹ See [REDACTED].

173 [REDACTED].

[REDACTED].¹⁷⁵ Thus, Euclid took advantage of the holes in the compliance programs for both distributors over that time period to obtain suspiciously high amounts of opioids that was not confirmed to be legitimate by either distributor.

B. CVS Store 3322

CVS Store 3322, located on Brookpark Road in Cleveland (Cuyahoga County), Ohio, ordered and received high-volumes of oxycodone from Cardinal Health and hydrocodone from CVS' own distribution centers from 2006 to 2014.¹⁷⁶

In the case of oxycodone, CVS 3322's volume began at █████ dosage units in █████ (an average of █████ per month, to an all-time high of █████ dosage units in █████ (an average of █████ per month).¹⁷⁷ The █████ level was only slightly reduced to █████ dosage units in █████ (an average of █████ per month), and still exceeded the █████ levels by more than █████.¹⁷⁸

Similarly, for hydrocodone, CVS 3322's volume began at [REDACTED] dosage units in [REDACTED] (an average of [REDACTED] per month) to a maximum of [REDACTED] dosage units in [REDACTED] (an average of [REDACTED] dosage units per month). However, CVS 3322 was [REDACTED] [REDACTED] [REDACTED]

[REDACTED]¹⁷⁹

Despite these high volumes of opioid purchases, neither Cardinal nor CVS appears to have adequately monitored this store or conducted appropriate investigations. The limited due diligence file produced by Cardinal begins in 2012 with an extremely limited due diligence memorandum located in the file.¹⁸⁰ Beyond the basic facts and figures in the file, Cardinal conducted several “site surveillance visits” from 2014 to 2017.¹⁸¹ These site surveillance visits were not the same as onsite visits and consisted primarily of observing the pharmacy from afar (e.g., looking for out-of-state license plates).¹⁸² Cardinal also did a single one-hour site

¹⁷⁵ See *infra* at Appendix A, Figure 2.

¹⁷⁶ See Memorandum from D. Roberts to File, OHIO CVS STORES LLC #3322 2007 BROOKPARK RD, CLEVELAND, OH 44109, 2 (Jul. 15, 2016) (Noting Cardinal was supplying all the store's Schedule II controlled substances needs and was the back-up distributor for Schedule III-IV controlled substances.), CAH MDL2804 00000216.

¹⁷⁷ See Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated Mar 25, 2019 at 1944-1945 (Opioid Shipments to AR7531418 by Distributor (2006-2014)).

178 *Id.*

¹⁷⁹ *See id.*

¹⁸⁰ See Memorandum from N. Rausch to File, CVS #3322 (DEA # AR7531418), (Jul. 17, 2012), CAH_MDL2804_00000204. However, there are records showing that CVS 3322 was purchasing from Cardinal as early as July 2007. See Cardinal Health Compliance Group, *Ingredient Limit Report*, (undated) (Showing orders in July 2007), CAH_MDL_PRIORPROD_DEA07_01120515 at 011208813-011208814.

¹⁸¹ See Cardinal Health, Surveillance Site Visit Report, (Jan. 24, 2014), CAH_MDL2804_00000207; Cardinal Health, Surveillance Site Visit Report, (Apr. 13, 2015), CAH_MDL2804_00000212; Cardinal Health, Surveillance Site Visit Report, (May 16, 2017), CAH_MDL2804_00000215.

¹⁸² See Stephen Forst Deposition, 30:4-33:19 (Jan. 22, 2019) (Mr. Forst was a Director in the Quality and Regulatory Affairs Department with anti-diversion responsibilities).

visit to the store in June 2012.¹⁸³ Neither the 2012 site visit nor the subsequent site surveillance visits noted any “red flags.” However, the way the visits were conducted (e.g., short duration or scope) they were of limited utility to detect diversion.

CVS, on the other hand, does not appear to have performed due diligence with respect to the high volume of HCPs shipped to this store, until May 2014.¹⁸⁴ Therefore, because neither Cardinal nor CVS performed adequate due diligence on the store, the pharmacy was able to purchase large volumes opioids uninterrupted from 2006 to 2014.

C. CVS Store 4800

CVS Store 4800 located at 590 East Market Street in Akron (Summit County), Ohio also ordered and received high-volumes of oxycodone from Cardinal Health and hydrocodone from CVS’ own distribution centers from 2006 to 2014.¹⁸⁵

CVS 4800’s oxycodone received from Cardinal Health ranged in volume from [REDACTED] dosage units in [REDACTED] (a monthly average of [REDACTED]) to a high of [REDACTED] dosage units in [REDACTED] (a monthly average of [REDACTED]).¹⁸⁶ For hydrocodone, the volumes ranged from [REDACTED] dosage units in [REDACTED] (a monthly average of [REDACTED]) to a high of [REDACTED] dosage units in [REDACTED] (a monthly average of [REDACTED]).¹⁸⁷ However, CVS 4800 [REDACTED]
[REDACTED] [REDACTED] [REDACTED].¹⁸⁸

The Cardinal due diligence file for CVS 4800 contains an undated questionnaire. Cardinal also conducted a single one-hour site visit to the store in June 2012¹⁸⁹ and two surveillance site visits in January 2014.¹⁹⁰ However, none of the visits noted any “red flags.” Furthermore, as in the case of CVS 3322, the way all three visits were conducted (e.g., short duration or scope) they were of limited utility to detect diversion. In the case of CVS, although this store did flag on the Item Review Reports, there was no documentation found demonstrating that diligence was performed by CVS with respect to these high-volume purchases.¹⁹¹ Therefore, because neither Cardinal nor CVS performed adequate due diligence on the store, the pharmacy was able to purchase large volumes opioids uninterrupted from 2006 to 2014

¹⁸³ See Cardinal Health, Report of Investigation, (Jun. 11, 2012), CAH, MDL2804_00000205.

¹⁸⁴ See CVS Health, Orders of Interest SOM-3724010, (May 5, 2014), CVS-MDLT1-000007490.

¹⁸⁵ See Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated Mar. 25, 2019 at 3216-3217. (Opioid Shipments to BR0287234 by Distributor (2006-2014)).

¹⁸⁶ *Id.*

¹⁸⁷ *Id.*

¹⁸⁸ *Id.*

¹⁸⁹ See Cardinal Health, Report of Investigation, (Jun. 11, 2012), CAH, MDL2804_00000688. Both site visits for CVS 3322 and 4800 were conducted on the same day.

¹⁹⁰ Cardinal Health, Surveillance Site Visit Report, (Jan. 20, 2014), CAH_MDL2804_00000692; Cardinal Health, Surveillance Site Visit Report, (Jan. 24, 2014), CAH_MDL2804_00000690;

¹⁹¹ See CVS, Item Review Report, (Jan. 4, 2011), CVS-MDLT1-000101092; CVS, Item Review Report, (Mar. 1, 2011), CVS-MDLT1-000101174; CVS, Item Review Report, (Mar. 29, 2011), CVS-MDLT-000101371.

PART V: Individual Company Reviews



9 McKesson Corporation

9.1 Background

Founded in 1833, McKesson Corporation (“McKesson”) describes itself as “the oldest and largest healthcare company in the nation, serving more than 50% of U.S. hospitals and 20% of physicians.”¹⁹² The company also claims to “deliver one-third of all medications used daily in North America.”¹⁹³ Within the U.S., McKesson currently operates 27 distribution centers registered with the DEA to handle controlled substances.¹⁹⁴ From 2009 to 2018, the number of McKesson employees grew from some 32,500 employees to 78,000.¹⁹⁵

McKesson’s controlled substances program can be characterized as having four distinct phases, which also correspond to the three different SOM programs McKesson employed. The first phase covers the period before May 2007, when McKesson’s program operated under Section 55 of the Drug Operations Manual. The second phase covers the introduction of the Lifestyle Drug Monitoring Program (“LDMP”). The third phase covers the

¹⁹² See MCKESSON CORPORATION, <https://www.mckesson.com/> (last accessed Nov. 27, 2018).

¹⁹³ *Id.*

¹⁹⁴ McKesson Corporation, *Board of Directors’ Response to International Brotherhood of Teamsters*, 1, P1.082 (April 20, 2018) at <https://www.mckesson.com/about-mckesson/newsroom/press-releases/2018/mckesson-board-releases-findings-and-recommendations-of-independent-investigation/> [MCK Teamsters Response].

¹⁹⁵ See STATISA, *Number of employees at the McKesson Corporation from 2009 to 2018*, <https://www.statista.com/statistics/240011/employment-at-the-mckesson-corporation/> (last accessed Dec. 6, 2018).

Controlled Substances Monitoring Program (“CSMP”), the successor to the LDMP, which was implemented as a result of the 2008 settlement with the DEA and periodically revised during the time period from 2008 to 2017.

With the 2017 settlement and the detailed obligations in the Compliance Addendum, McKesson’s controlled substances compliance program entered a new fourth phase.

9.2 Executive Summary

My review of McKesson’s controlled substances compliance efforts from 1996 to the present reveals a consistent pattern of systemic failures to meet its regulatory and corporate governance obligations with respect to controlled substances. Poor design exacerbated by poor implementation resulted in a series of “paper programs” that were neither effective for controlling diversion nor in identifying and reporting suspicious orders of controlled substances.

The root cause of this systemic failure largely comes down to corporate culture. McKesson’s controlled substances compliance efforts did not have the support and commitment of McKesson’s senior management and its Board of Directors. Despite the well-documented existence of the applicable compliance program standards for controlled substances and corporate compliance dating back to the 1970s and 1990s respectively, which McKesson management was aware of, McKesson did not attempt to implement effective compliance programs for either controlled substances or corporate compliance until many years later.

Furthermore, McKesson has twice settled with the DEA for the company’s inability to “maintain ... effective controls against diversion of particular controlled substances,” and its failure to “design and operate a system to disclose to the registrant suspicious orders of controlled substances.”¹⁹⁶ These settlements occurred in spite of repeated signals to McKesson and its management of the company’s obligations with regards to the CSA and acknowledgements by those in the position to judge “what good looked like” that its controlled substances compliance program was inadequate. McKesson’s management simply did not act unless it was pushed to do so by the DEA, and even then, the company only did the minimum to avoid further scrutiny.

9.3 Impact

The impact of McKesson’s failure to maintain an effective compliance program can be seen in Summit and Cuyahoga County. Based on an analysis done using McKesson’s own data, during the period from 2006 through mid-2008, no suspicious orders in either county were stopped or reported to DEA by McKesson.¹⁹⁷ This is in stark contrast to mid-2008 through the end of July 2013 when McKesson flagged [REDACTED] threshold excursions for pharmacies in Summit and Cuyahoga counties respectively.¹⁹⁸ Even so, McKesson

¹⁹⁶ See Administrative Memorandum of Agreement between United States Department of Justice, Drug Enforcement Administration and McKesson Corporation at 2 (Jan. 17, 2017), MCKMDL00355349 [“2017 AMOA”].

¹⁹⁷ See Analysis of Cuyahoga and Summit County Threshold Breaches & DEA SOM Reports from January 1, 2006 to July 31, 2013, MCKMDL00478912.

¹⁹⁸ *Id.*

made no suspicious order reports to the DEA involving any of [REDACTED] flagged orders in those two counties. .¹⁹⁹ This is despite the fact that in 2015, while negotiating with the DEA and moving towards its second settlement, McKesson reported on average 6.6 suspicious orders per customer to the DEA during that single year alone.²⁰⁰

As a result, various retail pharmacies obtained high levels of opioids with little or no investigation or interrogation. Below are a few illustrative examples.

██████████ was located in Summit County, Ohio and in 2015 was one of 17 pharmacies owned and operated by ██████████, a local grocery store chain.²⁰¹ ██████████

_____ . That request alone should have triggered a suspicious order report, but it did not.²⁰⁷

199 *Id.*

200

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

208 [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

The New Castle Distribution Center

McKesson has identified the New Castle Distribution Center as its facility that shipped more than 99% of the opioid products to retail pharmacy customers in both Summit and Cuyahoga Counties.²¹³ Therefore, this Distribution Center's due diligence practices regarding suspicious orders, are particularly relevant to my analysis. Below, I discuss in detail two specific pharmacies serviced by the New Castle Distribution Center.

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

213 [REDACTED]

216 *Id.*

217 [REDACTED]

220 *Id.*

[REDACTED]
 [REDACTED]
 [REDACTED]

221 [REDACTED]

[REDACTED]

[REDACTED]
 [REDACTED]
 [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED] [REDACTED]

[REDACTED]

9.4 Company Commitment – Compliance Culture, Organization & Resources

A horizontal bar chart showing the percentage of respondents who believe the U.S. should take action to address climate change, broken down by age group. The x-axis represents the percentage, ranging from 0 to 100. The y-axis lists age groups. The bars are black, and the chart includes a legend indicating that the bars represent 'U.S. should take action to address climate change'.

Age Group	Percentage (%)
18-29	92
30-49	85
50-69	90
70+	88
18-29	95
30-49	82
50-69	93
70+	96
18-29	94
30-49	80
50-69	91
70+	97
18-29	88
30-49	75
50-69	85
70+	90

9.4.1 McKesson's culture demonstrates a lack of commitment & support for complying with its controlled substances obligations.

Throughout the review period, McKesson repeatedly demonstrated that notwithstanding its assurances to the DEA that it took controlled substances compliance seriously, the opposite was true. In other words, McKesson failed “to walk the talk.” Even McKesson’s establishment of a Corporate Compliance program in the 2006 timeframe with the elevation of the General Counsel, Laureen Seeger to the role of Chief Compliance Officer, Executive Vice President, and General Counsel did little to change McKesson’s behavior.²³⁹ As outlined below, the Corporate Compliance function had little involvement with controlled substances compliance until the mid-2015 organizational changes. Even then the need to comply with the controlled sub[REDACTED] requirements does not appear on the Corporate Compliance “radar screen” (see discussion below).

This lack of commitment and support, and the resulting poor culture of compliance regarding controlled substances can be seen in the multiple warnings McKesson received during the period from the DEA, as well as the two DEA settlements that occurred within ten years of each other.²⁴⁰ Furthermore, prior to concluding each of its two government settlements, McKesson and the DEA had repeated contact, conversations and correspondence about the poor state of McKesson’s controlled substances program.²⁴¹ Therefore, McKesson was well aware of the DEA’s expectations with regards to anti-diversion and SOM programs, knowledge it shared with its employees on several occasions during the period.²⁴² However, McKesson failed to translate that awareness to fashion an effective controlled substances compliance program.

²³⁹ See Laureen E. Seeger, Executive Profile, BLOOMBERG, <https://www.bloomberg.com/research/stocks/people/person.asp?personId=26521629&privcapId=92001> (last accessed Dec. 21, 2018); see also Michael Johnsen, McKesson names new general counsel and chief compliance officer, DRUGSTORE NEWS (Jun. 23, 2014) at <https://www.drugstorenews.com/news/mckesson-names-new-general-counsel-and-chief-compliance-officer/>. However, as discussed in the Addendum, McKesson was a late bloomer when it comes to corporate compliance, as many others in pharmaceutical industry had established their programs years earlier and some were even on their second-generation program.

²⁴⁰ See Settlement and Release Agreement and Administrative Memorandum of Agreement between the U.S. Dept. of Justice, Drug Enforcement Administration and McKesson Corp. (May 2, 2008), MCKMDL00337001 [“2008 MOA”]; see also 2017 AMOA.

²⁴¹ See Memorandum from M. Mapes to J. Rannazzisi, *Internet Presentation with McKesson Corp. on September 1, 2005*, (Oct. 20, 2005), MCKMDL00496859; see also Memorandum from M. Mapes to J. Rannazzisi, *Meeting Between Office of Diversion Control (OD) and McKesson Corp. on January 3, 2006*, (Jan. 23, 2006), MCKMDL00496876; *Summary of DEA-HDMA Mtg. on Suspicious Orders*, (Sept. 7, 2007), MCKMDL00574906; Letter from W. Ihlenfeld, II to G. Hobart, *Claims Against McKesson Corporation*, (Nov. 6, 2013), MCKMDL00409048; Letter from D. Cutteman, *et al.*, to G. Hobart, *Registration Consequences for McKesson Corporation for Violations of the Controlled Substances Act*, (Nov. 4, 2014), MCKMDL00409453; Letter from W. Ihlenfeld, II to G. Hobart, *McKesson Matter – Ongoing Settlement Discussions*, (Mar. 20, 2014), MCKMDL00409174; Letter from J.F. Walsh to G. Hobart, *Possible Civil Action Against McKesson Corporation for Violations of the Controlled Substances Act*, (Aug. 13, 2014), MCKMDL00409224 at MCKMDL00409234; see also DEA 9/27/2006, 2/7/2007, 12/27/2007, 6/12/2012 Letters.

²⁴² See Presentation by D. Walker, *Lifestyle Drugs & Internet Pharmacies*, McKesson National Operations Conf., 4 (2007) (DEA Focus Slide lists what “DEA expects”), MCKMDL00403340 (Also stamped MCK-WVAG-003-0001335) [Walker Internet Presentation]; Denver Sales Mtg. presentation, *Controlled Substances Monitoring Program*, 5 (Mar. 10, 2008) (“Regulation *has not* changed, but the extent to which we are now required to monitor and provide stronger safeguards to ensure legitimate use of controlled substances *has*.” (Emphasis in the original)), MCKMDL00267635 at MCKMDL00267640 [“Denver Sales Meeting Presentation”]; see generally, Presentation by G. Boggs, *State of Prescription Drug Abuse*, (2013), MCKMDL00336833 [“Boggs Presentation”].

For example, in 2008 during the initial roll-out of McKesson's new Controlled Substances Monitoring Program ("CSMP"), the pharmacy program guide contained this message:

McKesson values you and your business and is committed to working closely with you to ensure that your pharmacy continues to be successful. This program addresses the DEA's requirements to ensure controlled substances are used in the way they were intended, but **it also ensures that you as a McKesson customer can continue with business as usual.** (Emphasis added).²⁴³

Since the CSMP is ostensibly a program intended to control the inappropriate procurement of controlled substances, the statement is self-contradictory. A controlled substances compliance program, which meets the DEA's requirements, cannot allow every customer "to continue with business as usual." There will be some "suspicious orders" that will remain "suspicious" even after a complete investigation; orders that will go unfilled and be reported to the DEA in an effective program.

McKesson, however, went so far as to even instruct its employees in the CSMP manual to avoid using the word "suspicious" because "[o]nce McKesson deems an order and/or customer suspicious, McKesson is required to act. This means all controlled substances sales to that customer must cease, and the DEA must be notified."²⁴⁴ These statements demonstrate a cultural disconnect between McKesson's actual culture and a good compliance culture. This cultural disconnect was something the government too observed and commented upon in its correspondence with McKesson.²⁴⁵

After the 2008 government settlement and fine, this cultural disconnect did not improve. For example, it can be seen in the 2013 presentation made to McKesson management by Gary Boggs, former Special Agent, and Executive Assistant to the Deputy Assistant Administrator for the DEA's Office of Diversion Control. Mr. Boggs made this presentation prior to McKesson hiring him in November 2013 as a Senior Director of Regulatory Affairs.²⁴⁶

During his presentation, Mr. Boggs shared with those present what he termed "recurring comments" concerning controlled substances compliance apparently made to him during his tenure with the DEA. The comments shared with McKesson's management fall into two different but intertwined categories. First, there are the comments that appear to shift the blame for the registrant's non-compliant situation to the DEA because the DEA:

- Won't talk to us;
- Won't tell us what to do;

²⁴³ See Email attachment from D. Walker to Bill de Gutierrez-Mahoney, *McKesson Controlled Substances Monitoring Program – Program Guide for Pharmacies*, attachment p. 2 (Apr. 4, 2008), MCKMDL00543610.

²⁴⁴ See McKesson, *McKesson Operations Manual for Pharma Distribution, Controlled Substances Monitoring Program*, 19 (Aug. 24, 2011), MCKMDL00000021 at MCKMDL00000039 ["CSMP Manual 2011"];

²⁴⁵ See J.F. Walsh 8/13/14 letter to G. Hobart at 11 ("Our investigation has revealed a disturbing pattern. McKesson-Aurora's desire for increased sales and retaining its customers overrode its obligations to report suspicious orders. We have identified this trend across several different areas."), MCKMDL00409224 at MCKMDL00409234.

²⁴⁶ See generally Boggs Presentation. Mr. Boggs apparently made this presentation after leaving the DEA and before being hired by McKesson, although according to William De Gutierrez-Mahoney, Boggs had committed to joining McKesson. See W. De Gutierrez-Mahoney Deposition, 63:6-13 (Nov. 28, 2018).

- Won't tell us what to look for;
- Won't tell us who the suspicious registrants are;
- Won't approve our SOM system; and
- Won't share ARCOS data with us.²⁴⁷

Second, there are a series of statements, such as:

- The customer's registration is still valid so...;
- Why couldn't we have been warned first;
- We're too busy to look at everything; and
- If we don't sell it, our competitors will,

These statements demonstrate, at best, a lack of knowledge about or at worst a disregard for the need and importance of having strong internal controls governing the distribution of controlled substances.²⁴⁸ In the case of McKesson, the narrative about the DEA not providing the company with enough direction to create an effective compliance program persists and has even been adopted by McKesson's Board of Directors.²⁴⁹

Additionally, despite statements to the DEA about McKesson's commitment to compliance,²⁵⁰ McKesson neither adopted nor made enhancements to its controlled substances program voluntarily. Rather McKesson claimed to be surprised that significant outside pressures were being placed upon it by the DEA. As Executive Vice President and Group President, Paul Julien, to whom U.S. Pharma reported, wrote in early 2006 "I was surprised and very troubled that the DEA was considering issuing a show cause against that [the Lakeland, Florida] facility."²⁵¹

In the same vein, both the 2008 and 2017 DEA settlements involved substantially the same issues,²⁵² and despite being under a settlement agreement with specific obligations to address the issues outlined in the 2008 settlement, McKesson failed to undertake the necessary program improvements and corrective actions to prevent the situation from recurring.²⁵³ Rather, the company required the DEA in 2017 to spell out those necessary improvements in specific detail, which was memorialized in the mandated Compliance Addendum.²⁵⁴

²⁴⁷ See Boggs Presentation at 27.

²⁴⁸ *Id.*

²⁴⁹ See MCK Teamsters Response at 9 and 29.

²⁵⁰ See, e.g., Letter from P.C. Julian to J.T. Rannazzisi, 1 (Jan. 18, 2006) ("let me assure you and DEA that McKesson is committed to a compliance program that ensures the safe distribution of health care products, especially controlled substances.") MCKMDL00571360 at MCKMDL00571361.

²⁵¹ See *id.*

²⁵² See 2017 AMOA at 2 ¶¶ 6-7 and 3 ¶2.

²⁵³ *Id.* at 3 ¶2 ("McKesson acknowledges that, at various, times during the period from January 1, 2009 up through and including the Effective Date of this Agreement . . . it did not identify or report to DEA certain orders placed by certain pharmacies that should have been detected by McKesson as suspicious based on the guidance contained in the DEA letters [referring to the Rannazzisi letters]."); see also 2017 AMOA at 4 ¶¶ 3b-3f.

²⁵⁴ See generally, Compliance Addendum to Administrative Memorandum of Agreement between United States Department of Justice, Drug Enforcement Administration and McKesson Corporation at 2 (Jan. 17, 2017) MCKMDL00355416 ["MKC Compliance

McKesson's approach to compensating its sales force also is indicative of the company's lack of support for anti-diversion efforts. Although the sales force was assigned specific and important responsibilities under the CSMP, their compensation plan was not aligned with those expectations thereby undermining the effectiveness of the anti-diversion program. From the earliest days of the CSMP, sales representatives were assigned an integral role in McKesson's anti-diversion program. For example, the September 2008 version of the CSMP, tasked sales representatives with several important due diligence tasks including:

- Providing reasoning behind sales that prompted a level 1 review following a threshold excursion;²⁵⁵
- Introducing customers to the requirements of the CSMP;²⁵⁶
- Assisting in the completion of customer questionnaires, including collecting dispensing data and conducting on-site physical inspections;²⁵⁷ and
- Assisting in the completion of customer interviews.²⁵⁸

There are two facets of this misalignment. First, the compensation plan for McKesson sales representatives until mid-2012 included opioid sales in the product mix.²⁵⁹ While compensation plans based on product sales are not *per se* wrong, in McKesson's case because there was no countervailing incentives for reporting suspicious customer orders and activity and undertaking the CSMP duties assigned to them, McKesson put a system in place that clearly valued sales and revenue above compliance and corporate responsibility.

Thus, until 2012 McKesson's sales representatives had a direct financial incentive for their customers to purchase opioids and other controlled substances, but McKesson never offered any financial or performance incentives to the sales force for reporting suspicious customer activity as it related to controlled substances or their duties under the CSMP.²⁶⁰

All of this is consistent with the point raised to McKesson's outside counsel in 2014 by the U.S. Attorney for the District of Colorado, who wrote "[o]ur investigation has revealed a disturbing pattern. McKesson-Aurora's

Addendum"]. Although this analysis focused on controlled substances compliance, McKesson adopted a similar approach towards its corporate compliance obligations. Despite a clear mandate from the FSGs in 1991 onwards, as well as the OIG in 2003 and the ACA in 2010, McKesson only established its corporate compliance program in 2006.

²⁵⁵ See McKesson Operations Manual, *Controlled Substance Monitoring Program*, 7, § 2.2.2 (Sept. 6, 2008), MCKMDL00533239 at MCKMDL00533246.

²⁵⁶ *Id.* at 8, § 3.1.

²⁵⁷ *Id.* at 9 § 3.2.

²⁵⁸ *Id.* at 13 § 3.3.

²⁵⁹ See Presentation by McKesson Corporation to DEA, U.S. Pharmaceutical Controlled Substances Monitoring Program, 19 (Dec. 17, 2014), MCKMDL00409483.

²⁶⁰ See Gene Cavacini Deposition, 127-180 (Jan. 25, 2019) (Discussing sales compensation plans for FY 2007 to FY 2013 and the lack of compliance measures.).

desire for increased sales and retaining its customers overrode its obligations to report suspicious orders. We have identified this trend across several different areas.”²⁶¹

9.4.2 McKesson’s contention that the SOM requirements were too vague for it to design & operate an effective SOM program is specious.

While McKesson has repeatedly argued that the standards surrounding a SOM program were unclear because of incomplete guidance from the DEA, that position simply does not track with the information I reviewed.²⁶² Just by reviewing the DEA regulations and general guidance letters provided to all registrants during the period, it is possible to get a clear concept of what a successful SOM program should look like.²⁶³ However, in addition to the publicly available information, McKesson received direct, non-public feedback from the DEA over a period of years.²⁶⁴ Finally, as a member of the Healthcare Distribution Management Association (“HDMA”), McKesson participated in and had access to the Association’s voluntary industry guidelines, “Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances,” which also provides clear direction on what a credible SOM program should look like.²⁶⁵

McKesson also understood that SOM programs are not static, but instead are meant to evolve as circumstances change.²⁶⁶ Consequently, McKesson’s assertion that it did not have enough information to construct and operate an effective SOM program demonstrates the company’s lack of commitment to effective controlled substances compliance.

²⁶¹ See J.F. Walsh 8/13/14 letter to G. Hobart at 11.

²⁶² The D.C. Circuit recently rejected this argument in *Masters Pharmaceuticals, Inc. v. DEA*, 15-1335 (D.C. Cir. 2017).

²⁶³ McKesson, itself, recognizes the regulations outline those responsibilities. See, e.g., Presentation, *McKesson’s Controlled Substances Monitoring Program - Regulatory Affairs Training*, 25 (undated), MCKMDL00336532 at MCKMDL00336549 [“RA 2015 Training”]; but see N. Hartle Deposition, 17:20-23 and 18:1-6 (Aug. 1, 2018) (Establishing via metadata that the training deck discussed in the De Gutierrez-Mahoney deposition was created on December 31, 2015); see *infra* Report Section 3.2.5.

²⁶⁴ See Memorandum from M. Mapes to J. Rannazzisi, *Internet Presentation with McKesson Corp. on September 1, 2005*, (Oct. 20, 2005), MCKMDL00496859; see also Memorandum from M. Mapes to J. Rannazzisi, *Meeting Between Office of Diversion Control (OD) and McKesson Corp. on January 3, 2006*, (Jan. 23, 2006), MCKMDL00496876; *Summary of DEA-HDMA Mtg. on Suspicious Orders*, (Sept. 7, 2007), MCKMDL00574906; Letter from W. Ihlenfeld, II to G. Hobart, *Claims Against McKesson Corporation*, (Nov. 6, 2013), MCKMDL00409048; Letter from D. Cutteman, *et al.*, to G. Hobart, *Registration Consequences for McKesson Corporation for Violations of the Controlled Substances Act*, (Nov. 4, 2014), MCKMDL00409453; Letter from W. Ihlenfeld, II to G. Hobart, *McKesson Matter – Ongoing Settlement Discussions*, (Mar. 20, 2014), MCKMDL00409174; Letter from J.F. Walsh to G. Hobart, *Possible Civil Action Against McKesson Corporation for Violations of the Controlled Substances Act*, (Aug. 13, 2014), MCKMDL00409224 at MCKMDL00409234

²⁶⁵ See Letter from Wendy Goggin to John Gray (Oct. 17, 2008) (discussing HDMA’s voluntary industry guidelines, “Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances.”) WAGMDL00673706.

²⁶⁶ See, e.g., RA 2015 Training at 27.

9.4.3 McKesson's organizational design of the controlled substances program reflects a lack of support for controlled substances compliance efforts.

Until approximately four years ago, the controlled substances program within McKesson operated like a junior function within the company. Controlled substances compliance, including SOM, is the purview of U.S. Regulatory Affairs and Compliance (formerly known as U.S. Regulatory Affairs).²⁶⁷

It was not until mid-2015 while preparing for the eventual January 2017 settlement with the government that McKesson created a connection between those responsible for the controlled substances program and the "governing authority" of McKesson (e.g., the Board of Directors and Corporate Officers).²⁶⁸ In the 2018 Teamsters Response report by McKesson's Board of Directors, the Board highlighted an example of what occurred in 2011 and 2012 because of that disconnect:

Based on the Committee's investigation, it appears that neither the Company's Board nor Audit Committee were made aware at the time of the issues raised by the DEA in connection with the Ohio distribution center inspection or the new distribution center registration application.²⁶⁹

However, McKesson's remedy for this governance disconnect is only a partial solution. The new connection, as shown in Figure 3 below,²⁷⁰ is an indirect ("dotted line") reporting relationship between the Senior Vice President, Regulatory Affairs and Compliance ("SVP, RA&C") and the Senior Vice President Compliance, Regulatory & Ethics (formerly, the Senior Vice President, Global Compliance & Ethics²⁷¹), who in turn reports directly to the Chief Compliance Officer ("CCO"), a function that was established in 2006.²⁷²

²⁶⁷ See Presentation, *Controlled Substances Compliance Program* at P1.1743.9 (Nov. 1, 2013), MCKMDL00516748 ["Regional SOM Training"]. The slide states that regulatory is responsible for "overall management and administration of the CSMP program," as well as "oversight and audit of recordkeeping and reporting."

²⁶⁸ See McKesson, *ISMC Controlled Substances Monitoring Program Operating Manual*, Version 1.1 at 6, § 3.1 (Jun. 1, 2015) (Version 1.0 was completed May 21, 2015), MCKMDL00337198 ["ISMC CSMP Manual 2015"]; see also RNA Controlled Substances Monitoring Program Operating Manual, Version 1.0, 5-6 (Apr. 17, 2018), MCKMDL00337622 ["RNA CSMP Manual 2018"]

²⁶⁹ See McKesson Teamsters Response at 17.

²⁷⁰ See McKesson, *ISMC Controlled Substances Monitoring Program Operating Manual*, Version 1.3 at 6, § 3.1, Figure 2 (Jan. 6, 2017), MCKMDL00395206 ["ISMC CSMP Manual 2017"].

²⁷¹ See ISMC CSMP Manual 2015 at 6, § 3.1, Figure 2.

²⁷² See Michael Johnsen, McKesson names new general counsel and chief compliance officer, DRUGSTORE NEWS (Jun. 23, 2014) at <https://www.drugstorenews.com/news/mckesson-names-new-general-counsel-and-chief-compliance-officer/>

Figure 3: Corporate Controlled Substance Governance Overview



Prior to the 2015 changes, U.S. Regulatory Affairs was a function located under the umbrella of U.S. Pharma Distributor Operations, which the Senior Vice President, Donald Walker led from 1997 to September 2015.²⁷³ Distributor Operations in turn reported to the U.S. Pharma Chief Operating Officer, who reported to the President of U.S. Pharma, who in turn reported to the Group Vice President, who reported to McKesson's Chief Executive Officer.²⁷⁴

As a result, before the 2015 changes, U.S Regulatory Affairs was at least six levels down in the organization from the CEO and seven levels down from the Board of Directors depending on which function is considered the governing authority for FSGs purposes. In other words, U.S. Regulatory Affairs, and hence the controlled substances program, lacked direct Corporate senior management oversight or support.²⁷⁵

Furthermore, the changes in 2015 to create the "dotted line" relationship between the SVP RA&C and the SVP Compliance, Regulatory & Ethics, in my opinion, are form over substance. Regardless of the "dotted line" into the Corporate Compliance function, the SVP RA&C's primary allegiance is to the President, U.S. Pharma as a direct report of the President. Thus, the SVP RA&C from 2015 to the present has the same reporting

²⁷³ See McKesson Corporation, Presentation to the USAO, Northern D. W. Va. and DEA, 7 (Mar. 12, 2014) (Walker identified as SVP, Distributor Operations), MCKMDL00409116 ["USAO Presentation 2014"], see also Donald Walker Profile at <http://www.spoke.com/people/donald-walker-3e1429c09e597c10039327ad> (last accessed Dec. 11, 2018) (Walker listed as SVP, Distribution Support in 2013).

²⁷⁴ See Presentation, *U.S. Pharma Regulatory Affairs/Compliance, All-Hands Meeting 2016*, 9 (Oct. 24-26, 2016), MCKMDL00336634 ("RA All-Hands") and see ISMC CSMP Manual 2015 at 6, § 3.1, Figure 2.

²⁷⁵ Although no formal governance link between the controlled substances compliance program and senior corporate management existed prior to 2015, there is evidence demonstrating McKesson Corporate was not totally unaware of what was happening. For example, in January 2006, Paul C. Julian, Executive Vice President, Group President for McKesson wrote a lengthy response to the DEA concerning discussion between McKesson and the DEA over the Lakeland Florida Distribution Center. See P.C. Julian 1/18/2006 letter to J.T. Rannazzisi.

relationship Mr. Walker had from 1997 to 2015. In short, nothing changed. McKesson could have accomplished significant change by simply making the SVP RA&C a direct report of the CCO with a dotted line to the President of U.S. Pharma, giving the SOM staff a true measure of independence.

In a similar fashion, the creation of the Regulatory Operating Committee (“ROC”) and the Controlled Substance Compliance Program National Governance Committee (“NGC”) in 2014 appears to be more form than substance.²⁷⁶ According to the 2015 ISMC CSMP Manual, the NCG’s duties are to provide high-level oversight of the CSMP; propose and adopt changes to the program; assure that concerns and inquiries regarding the program are resolved in a timely manner; monitor drug diversion trends and the effectiveness of the program; review significant compliance risk areas; and ensure proper communication of significant compliance risks.²⁷⁷ However, in the 2017 ISMC CSMP Manual, those duties were truncated to “high-level oversight of US Pharma’s compliance with the CSMP, [and] US Pharma’s compliance with the Controlled Substances Act and its implementing regulations.”²⁷⁸ With the exception of the Law Department and Internal Audit, NGC representatives are all senior U.S. Pharma employees.²⁷⁹ In short, this is the same group of employees that was in charge of the SOM program from pre-2000 to 2014.

The ROC, on the other hand, “is responsible for program-wide decisions regarding the CSMP, implementation, and execution of CSMP enhancements, hiring and onboarding of the Regulatory Affairs team, and supporting the technology and work needs of the Regulatory Affairs team.”²⁸⁰ This committee consists of the SVP RA&C and his or her senior directors.²⁸¹

Although the McKesson Board of Directors publicly touted how the company “enhanced its governance oversight of the CSMP”²⁸² by establishing the NGC and the ROC, neither of these two committees ironically has a formal place on the official organizational chart in the CSMP Manual.²⁸³ Nor do they have a direct reporting responsibility to the “governing authority” of McKesson (e.g., the Board of Directors and Corporate Officers). This is unlike the CCO, who has a formal reporting relationship to the Board Audit Committee. Therefore, it is unclear whether these committees perform real oversight and force change or are simply informal group designations established to give the embattled U.S. Pharma management team more credibility and legitimacy.

McKesson’s organizational design amounts to a serious deficiency as noted by the OIG, because “for a compliance program to be effective, it must have the support and commitment of senior management and the

²⁷⁶ See MCK Teamsters Response at 23.

²⁷⁷ See ISMC CSMP Manual 2015 at 9; *see also* MCK Teamsters Response at 23.

²⁷⁸ See ISMC CSMP Manual 2017 at 8.

²⁷⁹ See ISMC CSMP Manual 2015 at 9; *see also* MCK Teamsters Response at 23.

²⁸⁰ MCK Teamsters Response at 24; *see also* ISMC CSMP Manual 2015 at 7-8.

²⁸¹ See ISMC CSMP Manual 2015 at 7.

²⁸² MCK Teamsters Response at 23.

²⁸³ See ISMC CSMP Manual 2017 at 6. The ROC and NDC also are not mentioned in McKesson’s Annual Reports for FY2014 – FY 2016.

company's governing body,"²⁸⁴ and the compliance function needs to be able "to effectuate change within the organization as necessary or appropriate and to exercise independent judgment."²⁸⁵ The positioning of the controlled substances program so low in the corporate hierarchy and without any direct avenue to either McKesson's corporate officers or the Board of Directors is not a reasonable design to ensure effective compliance. In addition, it is another clear indication that McKesson neither supported nor is committed to ensuring that the SOM process and by extension the entire controlled substances program is optimized to be effective.

9.4.4 McKesson failed to resource the controlled substances program appropriately.

Before 2007 and the first DEA settlement, McKesson told the government that its regulatory affairs team consisted of 3 FTEs (2 SVPs and 1 Director of Regulatory Affairs).²⁸⁶ From 2007 to 2012, McKesson expanded the team to 10 FTEs by adding another Director of Regulatory Affairs, and 6 Regional RA directors, one of whom was former DEA investigator.²⁸⁷

The years from 2012 to 2014 marked a period of headcount expansion for McKesson's controlled substances program. Between 2012 and the March 2014 presentation to the U.S. Attorney's Office, McKesson added 2 Senior Directors and 6 Directors plus 2 analysts and 10 regulatory affairs managers of which 5 were former DEA employees. According to McKesson, this brought the team's total headcount to 30 FTEs.²⁸⁸

As noted above, McKesson distributes controlled substances from numerous distribution centers ("DCs") nationwide. While the number of DCs fluctuated during the review period, in 2014 McKesson told the U.S. Attorney's Office and the DEA that the company had 28 customer facing distribution centers plus 2 redistribution hubs.²⁸⁹ These centers serviced approximately 25,000 pharmacies and "processed 1.2 million order lines per night."²⁹⁰ Analyzing those numbers specifically for controlled substances provides the additional insight that even after the headcount expansion that began in 2012 McKesson still only had one controlled substances Regulatory Affairs person for each distribution center and redistribution hub, who oversaw approximately 1,240 controlled substances orders per night involving on average 833 customers.²⁹¹

²⁸⁴ See OIG Pharma Guidance at 23731.

²⁸⁵ *Id.* at 23739.

²⁸⁶ See USAO Presentation 2014 at 7.

²⁸⁷ See USAO Presentation 2014 at 8.

²⁸⁸ *Id.* at 9.

²⁸⁹ See USAO Presentation 2014 at 4.

²⁹⁰ *Id.* Another significant headcount increase occurred once more when McKesson prepared to conclude its second settlement with the DEA that was consummated in January 2017. See ISMC CSMP Manual 2017 at 6, § 3.1, Figure 3.

²⁹¹ Using the data from Appendix C, Figure 1, the formula for number of nightly controlled substances orders was [1.2 million (number of daily line times) x .031 (% of controlled substances sales (using lowest figure))] /30 (number of distribution centers) = 1,240. Assuming every McKesson customer sells controlled substances in 2014 (25,000 customers)/30 FTEs = 833.33 customers per staff member.

Comparing this data with the number of full-time equivalents (“FTEs”) in the controlled substances program reveals two important insights. First, the increase from 3 FTEs to more than 40 over 10 years appears to be a tacit admission by McKesson that the company knew its SOM program was under-resourced.²⁹² Second, even with the headcount additions provided to the program by March 2014, McKesson’s controlled substances program remained significantly under-resourced when factoring in the DEA’s basic expectations to “know your customer” and report suspicious orders.

Substantial compliance headcount increases are a common response to settlement agreements as a company seeks to integrate the additional burden of administering the settlement provisions into its customary, pre-settlement workload. For example, as a result of their Corporate Integrity Agreements, both Pfizer and GSK significantly expanded their compliance teams. Compliance headcount increases also are commonplace as a normal part of organizational growth. During the review period, McKesson experienced significant revenue growth.²⁹³ However, these headcount increases due to organic growth are usually more incremental than exponential. In my experience, this dramatic an increase in compliance headcount is a clear indication that McKesson recognized its controlled substances program was significantly under-resourced.

[REDACTED]

Working with McKesson’s data disclosed to the U.S. Attorney’s Office and others, at the 2014 staffing level of 30 FTEs for SOM and assuming every team member is doing some suspicious order reviews, each team member had to review approximately 18 new suspicious orders per day.²⁹⁶ By 2017, that number drops to approximately nine (9) per day per staff member, which is still a significant workload.²⁹⁷

²⁹² See ISMC CSMP Manual 2017 at 6, § 3.1, Figure 3.

²⁹³ See MacroTrends, *McKesson Revenue 2006 to 2018* at <https://www.macrotrends.net/stocks/charts/MCK/mckesson/revenue> (last accessed Dec. 12, 2018) (Showing McKesson revenue more than doubled during the period).

²⁹⁴ See McKesson Operations Manual for Pharma Distribution, *Lifestyle Drug Monitoring Program*, 1 (May 16, 2007) (emphasis in the original) (This program was only in effect from May 2007 until May 2008 when it was replaced by the CSMP), MCKMDL00330211 [“LDMP Manual”]; CSMP Manual 2011 at, 19 (“McKesson’s responsibility is to “Know our Customer.”); see also Boggs Presentation at 39 (Sept. 30, 2013).

²⁹⁵ See McKesson CSMP “Red Flags,” 1 (May 2015), MCKMDL00335740; see also CSMP Manual 2011, § 6.1; see also *Masters Pharmaceutical, Inc. v. DEA*, No. 15-1335, Op. 5 (D.C. Cir. 2017) (“Once a distributor has reported a suspicious order, it must make one of two choices: decline to ship the order, or conduct some “due diligence” and—if it is able to determine that the order is not likely to be diverted into illegal channels—ship the order (the Shipping Requirement).” citing DEA, *Southwood Pharmaceuticals, Inc – Revocation of Registration*, 72 Fed. Reg. 36487, 36500 (Jul. 3, 2007)); See James Arnold presentation, *Effective Controls Against Diversion*, Manufacturer/Importer/Exporter Conference, 41 (Jun. 2013) available at https://www.deadiversion.usdoj.gov/mtgs/man_imp_exp/conf_2013/.

²⁹⁶ Using the data in Appendix C, Figure 1, the formula used was [3-year average of suspicious orders/number of days per year]/number of FTEs = result. Therefore, $((230,000 + 220,000 + 145,000)/3)/365/30 = 18.1$.

²⁹⁷ Again, from Appendix C, Figure 1 data, $[(145,000 \text{ number of SOs in 2017})/365 \text{ days per year}]/44 \text{ FTEs} = 9.02$.

On top of reviewing suspicious orders, the controlled substances staff members are each responsible for “knowing” approximately 833 customers, including performing onboarding, and actively monitoring the information that could affect ordering patterns (e.g., region served, county population, sale representative visit information, etc.).²⁹⁸ By 2017 that number had climbed to 909 customers per staff member.²⁹⁹

Finally, the controlled substances program staff members also have other compliance “back office” work such as training, writing reports, updating policies and procedures, etc. to address. The bottom line is that even with McKesson’s headcount expansions post-2014, the compliance workload per person is not sustainable to achieve effective compliance.

9.5 Program Core - Requirements, Education, Detection & Corrections

Written standards, employee education, detecting and correcting breaches of the company’s written standards and the law are at the core of an effective compliance program. However, in each of these areas, the data I reviewed demonstrates that McKesson came up short such that McKesson’s controlled substances program overall was ineffective to detect and prevent instances of controlled substances diversion during the review period. Furthermore, McKesson knew or should have known that its program was inadequate.

9.5.1 McKesson’s Code of Conduct does not meet generally accepted compliance standards

The earliest version of the McKesson Corporation Code of Business Conduct and Code of Ethics (“Code”) reviewed dates to February 2009.³⁰⁰ However, the Code’s copyright footer notes 2005, 2006 and 2009 suggesting that McKesson’s first Code dates to 2005 or 2006, approximately the same time that McKesson named its first Chief Compliance Officer.³⁰¹ The Code states that it “applies to all McKesson personnel, which includes every McKesson officer, director, employee, and agent.”³⁰²

The Code also contains a section entitled “Interacting With Government” that is focused primarily on federal health care fraud and abuse laws.³⁰³ While the Code covers many of nuances such as the Federal False Claims Act, the Foreign Corrupt Practices Act and lobbying, nowhere is controlled substances compliance highlighted. Thus, controlled substances compliance is at best covered under the general catch-all “we should ... [c]onduct all work and business affairs lawfully and with integrity.”³⁰⁴

²⁹⁸ This analysis assumes that every McKesson customer sells controlled substances. Therefore, using 2014 data from Appendix C, Figure 1, 25,000 customers/30 FTEs= 833.33 customers per staff member.

²⁹⁹ Using 2017 data from Appendix C, Figure 1, 40,000/44 = 909.09 customers per staff member.

³⁰⁰ See McKesson Corp., *McKesson Corporation Code of Business Conduct and Code of Ethics Code* (Feb. 2009), MCKMDL00375003 [“2009 Code”]

³⁰¹ *Id.*

³⁰² *Id.* at 1.

³⁰³ *Id.* at 20-26.

³⁰⁴ *Id.* at 1-2.

Since 2009, the McKesson Code has been amended several times, with the most recent changes being made in August 2018.³⁰⁵ However, while the Code of Conduct continues to specifically address certain standard risk areas found in most company codes, McKesson's 2018 Code still does not have a corresponding specific section addressing controlled substances compliance. This is despite McKesson entering into settlement agreements with the DEA on multiple occasions and paying substantial fines.³⁰⁶ The 2018 version also does not contain an "Interacting With Government" section.

The lack of any mention of complying with the controlled substances laws and regulations is at odds with McKesson's representations in the 2017 Compliance Addendum that:

McKesson understands that policies and procedures, such as a Code of Conduct, are central components to an effective controlled substance monitoring program, and further, that maintenance and oversight of policies to address DEA requirements create a culture of compliance in support of those requirements.³⁰⁷

Generally, a company's code of conduct addresses the compliance risk areas that are most important to the company, which is the point the Compliance Addendum is making. Therefore, by not including a section on controlled substances in its Code of Conduct, McKesson's does not meet generally accepted compliance standards and is not aligned with the Compliance Addendum's representations about McKesson's understanding of the Code's importance.

9.5.2 McKesson's standards outlining its controlled substances program were poorly organized and drafted undermining the primary purpose for creating standard policies and procedures.

Throughout the review period, McKesson's written standards outlining its controlled substances program were poorly organized and drafted which undermined the primary purpose of standard policies and procedures – achieving consistent compliance. Length, poorly defined objectives and ambiguous definitions all contribute to the program manuals being difficult to traverse and determine how the SOM process operates.

A. Drug Operations Manual, Section 55

The earliest manifestation of McKesson's controlled substances program that I reviewed is dated from 1997.³⁰⁸ Informally known as Section 55, this was the program manual in use from at least 1997 to May 2007. The

³⁰⁵ See McKesson Website, <https://www.mckesson.com/investors/corporate-governance/code-of-conduct/>, (last accessed November 29, 2018) (Amendments occurred in 2013, 2016 and 2018). The 2018 amendments occurred because McKesson "updated its Code of Conduct to reflect rebranding of its business operations in the European Economic Area and European Union under the consolidated name McKesson Europe, including translated versions for each of the different sub-brands and countries in which the company operates."

³⁰⁶ See McKesson Code of Conduct, Table of Contents at ii, available at <https://www.mckesson.com/documents/investors/mckesson-code-of-conduct/> (last accessed Dec. 5, 2018). The Code does contain the standard compliance statement that "[w]e comply with applicable laws everywhere we do business around the world." *Id.* at 1.

³⁰⁷ See MCK Compliance Addendum at 2, section § II.A.

³⁰⁸ See Memorandum from D. White to Distribution, *Drug Operations Manual Section 55-DEA Compliance* (Jan. 15, 1997) (MCKMDL00651873).

controlled substances chapter is a 137-page, detailed recitation of the DEA regulations, and a description of how to fill out various required forms, physical security requirements, which also includes a section on suspicious orders.³⁰⁹ Section 55 is missing standard procedural sections such as responsibilities and definitions. The chapter also does not contain step-by-step instructions to tell an employee how to perform suspicious order monitoring. Therefore, as a basic standard operating procedure, Section 55 is deficient.

Section 55 also does not meet the basic DEA requirements for a SOM Program. In the suspicious order section (IV), Section 55 outlines five different reports concerning a customer's purchases (Controlled Substances Sales Report, Controlled Substances Customer Purchase Report, Daily Controlled Substance Suspicious Order Warning Report, Monthly Controlled Substance Suspicious Purchases Report and the Monthly ARCOS Customer Recap Variance).³¹⁰ Despite the fact that the section quotes the DEA requirements around suspicious orders, and lists these system generated reports, there was no apparent requirement for McKesson employees, Distribution Center Managers, Operations Managers or Warehouse Staff (DCM/OM/WS) to take any action other than review the Suspicious Order Warning Reports, sign to acknowledge the review had been done nightly, fax a copy to the local DEA Office and file it in the appropriate file folder.³¹¹ Thus, there was no documented obligation for the McKesson staff members to place the orders on hold or investigate the reason for the customer appearing on the report.

The output from these reports, known as DU-45 reports, was also rudimentary. Customers appeared on the DU-45 report when sales of controlled substances, including opioids, to that customer exceeded three times of that customer's 12-month purchase average for that base code.³¹²

These standards are clearly at odds with the DEA requirements that distributors must know their customers and therefore must investigate the circumstances surrounding their orders to determine if any should be submitted to the DEA as suspicious and/or whether the order appears to be indicative of diversion and stopped.³¹³ As outlined by Dave Gustin, "the previous reports [under Section 55] were not the exclusive and proper response to this regulation. We have an obligation to report 'suspicious orders.' With no clear definition of what constitutes a suspicious order, we must rely on our own judgment as to what that is. If we report anything, we believe to be truly suspicious we will be meeting the spirit and letter of the regulation. Simply reporting larger than usual orders does not when there are so many plausible and routine reasons for orders to be 'larger than normal.'"³¹⁴

³⁰⁹ See generally McKesson, *Drug Operations Manual Section 55 DEA Compliance* (Jul. 2000), MCKMDL00346554) ["Section 55"].

³¹⁰ See Section 55 at 28-30, § IV.A.2.a-e.

³¹¹ See Section 55 at 31-32, § IV.B.2 (Section IV.B.2.d also notes that "[w]e are required by federal law to report suspicious orders upon discovery.").

³¹² See McKesson, *Drug Operations Manual Section 55 DEA Compliance*, MCKMDL00651919-20 (Jan. 27, 1997), MCKMDL00651873; Gary Hilliard Deposition, 163:21-169:7 (Jan. 10, 2019).

³¹³ See Discussion *infra* at Section 5.3.

³¹⁴ See Email from D. Gustin to T. Williams, *et al.*, RE: Variance and Suspicious Reports (Feb. 4, 2011), MCKMDL00510747; see also McKesson, DU45R05B8165 Monthly Controlled Substance Rpt., (Apr. 3, 2007) (This example report is 647 pages), MCKMDL00660789

B. Lifestyle Drug Monitoring Program

Although extremely short-lived (May 2007 to May 2008), McKesson's next documented program was the Lifestyle Drug Monitoring Program or "LDMP." The program was created as a result of the DEA's concerns over internet pharmacy sales of controlled substances.³¹⁵ Although considered by McKesson as a controlled substances compliance program, the LDMP program does not constitute a complete controlled substance compliance program as it was limited to just four drug products (oxycodone, hydrocodone, alprazolam, and phentermine).³¹⁶ However, during the period the LDMP existed, McKesson distributed other controlled substances such as hydromorphone, methadone, and morphine,³¹⁷ but McKesson did not monitor those other drugs. Instead, McKesson only monitored what the DEA "told them to" and expressly noted that "[t]he list of substances monitored by McKesson will only be adjusted when and if the DEA focus list is modified."³¹⁸ Thus, McKesson committed to only doing the bare minimum, but from a regulatory perspective that "bare minimum" was deficient from the outset.

Although the LDMP manual is closer to a conventional SOP format, it still lacks critical components such as definitions, an outline of the specific responsibilities for key personnel (responsibilities section), effective date and revision history.³¹⁹ As a result, it is impossible for an employee to know (other than taking it at face value) whether the document was properly approved, or when it became effective. These deficiencies create a large compliance loophole. Nor is it clear whether general employees at McKesson's distribution centers have any responsibilities under the LDMP or whether that is reserved for employees at the level of DCM and above.

C. Controlled Substances Monitoring Program

Replacing the LDMP, the Controlled Substances Monitoring Program or "CSMP" came on-line in 2008, and although modified on several occasions, it remains in use today. A review of the 2011 Manual, which is substantially similar to the 2008 version for this discussion, reveals that McKesson added a revision history section to the manual.³²⁰ As with the previous program manuals, the 2011 CSMP still lacks a formal responsibilities section, which makes it difficult for various functions and employees in general to determine their roles.

With the 2015 introduction of two CSMP manuals, one for the Independent Small Medium Chains ("ISMC")³²¹ and ultimately one for Retail National Accounts ("RNA") customers,³²² McKesson finally adopted the standard

³¹⁵ See Walker Internet Presentation at 3-6. *see also* See, e.g., Sandy Campbell, *Lifestyle Drug Program*, McKesson U.S. Pharma DEA Licensure Audit, 1 (last revised Jul. 27, 2007), MCKMDL00591949-00591953 ["LDMP Narrative"]; Letter from J. Gilbert to L. Barber (Apr. 25, 2007) (Mr. Gilbert from Hyman, Phelps & McNamara was McKesson's outside counsel and Mr. Barber was Associate Chief Counsel in the Office of Chief Counsel, Diversion and Regulatory Litigation Section, DEA), MCKMDL00330924 (Also stamped MCK-HOI-002-0000001).

³¹⁶ LDMP Narrative at 2.

³¹⁷ See McKesson Regional Statistical Norms, 2 (Feb. 24, 2014), MCKMDL00430387 at MCKMDL00430388.

³¹⁸ See LDMP Narrative at 2.

³¹⁹ See LDMP Manual.

³²⁰ See CSMP Manual 2011, 24-26.

³²¹ See ISMC CSMP Manual 2015.

policy and procedure elements, but at the same time it increased the program's complexity by creating two separate CSMP manuals. These dual manuals necessitate the use of additional resources to ensure the CSMP provisions common between them are always aligned, and potentially raising questions (and compliance breakdowns) for employees as to which manual governs their day-to-day activities, as distribution centers service both ISMC and RNA customers. This added complexity was something that McKesson recognized, but nevertheless persisted in keeping both manuals.³²³

9.5.3 McKesson's *ad hoc* approach to controlled substances education for its employees did not and still does meet generally accepted compliance standards.

Although an effective compliance program is expected to have a formal and robust training and education program surrounding the requirements for a controlled substances distributor and how McKesson meets those requirements, there is scant evidence that McKesson had any kind of a systematic approach to controlled substances training prior to 2015. In fact, two LDMP audits conducted in 2007 noted that "[t]he greatest barrier to success of the LDMP is the lack of understanding of the DEA expectations."³²⁴ This was also aptly illustrated by Blaine Snider in his deposition when he testified that although he followed the Code of Federal Regulations pertaining to the distribution of controlled substances and SOM, he did not know the phrase "legitimate medical purpose" nor had he heard of the Controlled Substances Act or the September 2006 DEA letter to all registrants.³²⁵

Section 55, the LDMP Manual and prior iterations of the CSMP Manual all fail to reference a training program.³²⁶ [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

³²² See RNA CSMP Manual 2018. While started in 2015 when the separation occurred (*see* Email from A. Palmer to M. Oriente, Family Code Request Examples, (Aug. 20, 2015), MCKMDL00628803.), the process was not completed until April 2018 more than 2-1/2 years later during which time the RNA group operated under SOPs that technically did not apply to them.

³²³ See Email from L. Brenner to N. Hartle, FW: RNA Operating Manual – Next Steps, (Apr. 11, 2018) ("The key thing we have to keep in mind is that to the extent this Manual is different from the ISMC Manual, the DEA will likely ask why, and we have to be prepared with defensible answers. So, one of the helpful things to discuss or flesh out in the RNA Ops Manual is why there is for example more flexibility in places, or less detail, or less mandatory specific documentation or diligence."), MCKMDL00570154.

³²⁴ See Gwen Allen, *Lifestyle Drug Program, McKesson U.S. Pharma – DEA Licensure Audit, So Cal DC*, 4 (Aug. 23, 2007), MCKMDL00591858; *see also* Sandy Campbell, *Lifestyle Drug Program, McKesson U.S. Pharma – DEA Licensure Audit, Washington Court House*, 3 (Aug. 10, 2007), MCKMDL00591862.

³²⁵ See Blaine Snider Deposition, 23:4-13, 24:4-15, 34:4-6 and 74:1-6 (Nov. 8, 2018). Snider was Director of Operations for the New Castle, PA distribution center from 2000 to the present. *Id.* at 19:7-13.

³²⁶ See generally Section 55; LDMP Manual; CSMP Manual 2011.

³²⁷ See McKesson Operations Manual, *Controlled Substances Monitoring Program*, 25, § 8 (Mar. 21, 2013), MCKMDL00002509 ["CSMP Manual 2013"].

[REDACTED]
[REDACTED]
[REDACTED] Sharon Longwell, Vice President of National Account Support Services, responded: “I have copied the [CSMP] manual so my team can see it because I don’t think we have ever seen this manual before.”³²⁹ This exchange between Mrs. Jonas and Ms. Longwell indicates that (a) routine controlled substances training did not occur prior to 2013, and (b) that McKesson failed to ensure that the CSMP manual was distributed to all employees needing access to it.

Also, I have seen no examples of standard on-boarding training materials before 2015. Furthermore, when the Regulatory Affairs group met on March 5-6, 2008 to discuss the roll-out of the original CSMP program, the meeting notes did not reflect any discussion about training or even list it as an item to be covered.³³⁰

[REDACTED]
[REDACTED]
[REDACTED]. Although this represents an improvement from the previous undocumented, *ad hoc*, approach, the standard controlled substances training was still significantly flawed, which has carried through to the current program.³³²

[REDACTED]
[REDACTED]
[REDACTED] This is a nebulous standard that lacks sufficient rigor and is widely variable.

³²⁸ See Email from T. Jonas to S. Longwell, *et al.*, Required Controlled Substance Training (May 1, 2013) (“p.s. Managers as identified in PeopleSoft are CC’d in this email.”), MCKMDL00521761 at MCKMDL00521763.

³²⁹ See Email from S. Longwell to T. Jonas, RE: Required Controlled Substance Training (May 2, 2013), MCKMDL00521761 at MCKMDL00521762.

³³⁰ See Email M. Oriente to D. Walker, *et. al*, Regulatory Meeting 3/5 & 3/6, Attachment (Mar. 7, 2008), MCKMDL00545048; *cf.* Email from C. Scofield to T. Jones, *et. al*, CSMP Sales Presentation (Mar. 7, 2008), MCKMDL00267635 (Training was not mentioned in the presentation either) [“CSMP Sales Presentation 2008”]; RA 2015 Training at 48.

³³¹ See ISMC CSMP Manual 2015 at 46, § 9.

³³² See ISMC CSMP Manual 2017 at 43 § 9; *cf.* ISMC CSMP Manual 2015 at 46, § 9. Apart from some minor changes, section 9 remains largely unchanged between the 2015 and 2017 versions.

³³³ See ISMC CSMP Manual 2015 at 46, § 9.1.

³³⁴ See ISMC CSMP Manual 2015 46, § 9.1.

³³⁵ See ISMC CSMP Manual 2015 46, § 9.2.

9.5.4 As early as 2005, McKesson knew its SOM program was not in compliance with DEA requirements.

In September 2005, the DEA met with McKesson representatives, during which the Agency highlighted two specific McKesson pharmacy customers with suspicious ordering patterns for hydrocodone.³³⁹ As part of the discussions, the DEA also reminded McKesson that it was required to: (a) report suspicious orders to DEA when discovered, (b) not rely on the DEA to determine if a suspicious order is legitimate, and (c) stop selling if they detected diversion.³⁴⁰ McKesson acknowledged in the meeting that it understood those obligations under the CSA.³⁴¹

over [REDACTED] period, to pharmacies previously identified by DEA to McKesson Corp.³⁴⁴ At the meeting, Gary Hillard, Director of Regulatory Affairs for McKesson, also disclosed to the DEA that McKesson's monitoring efforts for hydrocodone were deficient because they only tracked branded products and did not include generics.³⁴⁵

Individually, [REDACTED]; all in approximately one-third of a month. Clearly those purchases and purchasers were "suspicious" and warranted further investigation whether through a simple application of common sense or the DEA's recitation of circumstances that might indicate diversion.³⁴⁶ Furthermore, sales of that magnitude also should not have continued until cleared by thorough company investigations.³⁴⁷ Therefore, by the time McKesson rolled out the LDMP in 2007, the company already had customers engaging in diversionary behavior that the DEA repeatedly noted and brought to the company's attention, but which McKesson failed to address properly.

9.5.5 McKesson's LDMP continued the company's non-compliance due to poor program design and implementation.

While the LDMP can fairly be characterized as a rudimentary attempt at a controlled substances compliance program, the LDMP design and implementation flaws were such that they predictably resulted in customers exceeding the 8,000-dosage unit monthly thresholds. As a result, the LDMP cannot be legitimately viewed as providing "effective controls against diversion of particular controlled substances into other than legitimate medical, scientific, and industrial channels."³⁴⁸

A. Establishing the Thresholds

The LDMP Manual describes the monitoring program as follows:

[The] Daily Dosage Summary and Dosage Limit Tracking Detail have been developed to allow McKesson to monitor customer purchases of controlled substances by the net number of dosage units sold based on the DEA's Controlled Substance Generic Base Code ID. McKesson will investigate customer activity when sales of a given generic base ingredient exceed a predefined dosage unit threshold within a calendar month. ... The same dosage threshold will be used for all classes of customers.³⁴⁹

³⁴⁴ See *id.*

³⁴⁵ See *id.*

³⁴⁶ See DEA 9/27/2006 Letter at 2.

³⁴⁷ See, e.g., DEA 6/12/2012 Letter at 2.

³⁴⁸ See 21 U.S.C. § 823 (b)(1).

³⁴⁹ See LDMP Manual at 1.

The threshold level was set at a generic level of 8,000 by the Regulatory Department.³⁵⁰ While having a set threshold is an improvement over the previous version of the SOM program, McKesson does not appear to have either a logical or statistical basis for setting the level at 8,000 dosage units per month. The absence of a sound basis for the threshold level manifested itself in two ways.

First, the threshold level applied to all classes of customers from independent, small and medium pharmacies to large national pharmacy chains. Having a single, uniform threshold did not comport with the DEA's requirements to "know your customer" and the need to individualize diversion detection. Thus, it appears that McKesson designed the LDMP to "check the box" that it had a controlled substances compliance program, rather than to develop a truly effective process.

Second, McKesson was aware that the DEA believed that the threshold level triggering further investigation was 5,000 dosage units per month on average.³⁵¹ McKesson, however, set its threshold level to trigger further investigation a full 37.5% higher. Although there is no definitive documentation outlining why McKesson set the threshold so much higher than DEA expectations, Paul Julian's letter to the DEA in January 2006 provides a possible explanation. In his letter, Mr. Julian referenced the DEA's 5,000 dosage unit expectation and went on to note that McKesson had more than 85 pharmacy customers ordering more than 5,000 dosage units of hydrocodone per month from the Lakeland, Florida distribution center alone.³⁵² Therefore, one possible explanation is that McKesson set the threshold level at 8,000 dosage units per month to avoid the need to make potentially disruptive adjustments to the numerous customers that were ordering more than 5,000 dosage units per month of hydrocodone.

B. The Daily Dosage Summary Report

Everything in the LDMP program keyed off the Daily Dosage Summary report. However, as Sandy Campbell's narrative documenting the LDMP in July 2007 revealed, the Daily Dosage Summary report was flawed in at least two crucial respects. First, she noted that "it is possible not all of the products containing one of the generic ingredients are included."³⁵³ This failure to monitor generic products was the same failure Gary Hillard told the DEA about during their January 3, 2006 meeting more than 18 months earlier.³⁵⁴ It remained uncorrected in the latest iteration of McKesson's SOM system although according to Ms. Campbell's narrative it would be corrected at some unspecified future date.³⁵⁵ The second flaw she noted was that the Daily Dosage Summary report was organized by distribution center ("DC"), and therefore a customer could both exceed the monthly 8,000 dosage unit threshold and avoid detection by spreading its purchases across multiple distribution

³⁵⁰ See LDMP Manual at 2, § 1.1; LDMP Narrative at 2.

³⁵¹ See P.C. Julian 1/18/2006 letter to J.T. Rannazzisi, at 3-4 ("DEA has stated that monthly sales of over 5,000 dosage units of hydrocodone should be used as a flag to whether the pharmacy is dispensing legitimate prescriptions."); see also Walker Internet Presentation at 4

³⁵² See P.C. Julian 1/18/2006 letter to J.T. Rannazzisi at 4.

³⁵³ LDMP Narrative at 3.

³⁵⁴ See M. Mapes 1/23/2006 memorandum o J. Rannazzisi at 2.

³⁵⁵ See LDMP Narrative at 3.

centers.³⁵⁶ Therefore, the base report the Distribution Center Managers (“DCMs”) relied upon to identify potentially suspicious orders was corrupted rendering the program ineffective from the outset.

C. LDMP Escalation Process

The LDMP program provided for a 3-tier escalation process that was triggered once a customer exceeded the threshold of 8,000 dosage units in a month.³⁵⁷ Once triggered, the DCM was instructed by the LDMP manual to evaluate the customer’s previous three month’s purchases, and examine whether:

- Previous sales were validated and approved.
- Sales have not increased more than 25% from any previous month.
- Sales are not increasingly steadily.
- Sales are consistent with customer type.
- Sales are consistent with any previous Sales or Customer communication.³⁵⁸

Once complete, “if the evaluation indicates that the customer’s purchases are reasonable and that no further investigation is required,” the DCM could approve the shipment.³⁵⁹ Only if the evaluation was inconclusive was Level 2 invoked, and Distributor Operations or Regulatory Affair was notified, as well as a site visit and a customer interview initiated.³⁶⁰ In addition to the substantial amount of work required for each excursion above the threshold, the LDMP program contained no objective criteria for determining when purchases were reasonable or what constituted an inconclusive evaluation.

The escalation process also did not provide for a hard cut-off to customers being reviewed to prevent potential diversionary orders from being filled.³⁶¹ This is in direct contravention to DEA’s expectation and explicit directions as outlined in the DEA’s September 27, 2006 letter written before the start of the LDMP program that;

a distributor has a statutory responsibility to exercise due diligence to **avoid filling suspicious orders that might be diverted** into other than legitimate medical, scientific, and industrial channels. . . the distributor should exercise due care in **confirming the legitimacy of all orders prior to filling**.³⁶²

By not providing hard cut-offs, McKesson created a “paper program” (e.g., one that looks good on paper, but in practice does little to achieve compliance) and allowed potential diversionary situations to continue unabated. For example, there was the case of Franklin Pharmacy (also discussed separately above). In November 2007,

³⁵⁶ *Id.*

³⁵⁷ *See* LDMP Manual at 2, § 1.1

³⁵⁸ *Id.*

³⁵⁹ *Id.* at 3, §§ 1.2 and 1.4.

³⁶⁰ *Id.* at 3-4, §§ 2.1 to 2.4.

³⁶¹ *See* W. De Gutierrez-Mahoney Deposition, 584:11-17 (Admitting that there was no hard blocking of orders submitted by customers undergoing LDMP review).

³⁶² DEA 9/27/2006 Letter at 2 (emphasis added.)

[REDACTED]

In addition, as these examples illustrate, McKesson misrepresented how the program worked to the DEA. In April 2007, McKesson's outside counsel wrote to the DEA that the company would not ship more than 8,000 dosage units per month to customers until due diligence was completed and would terminate and notify DEA about any account "where it cannot adequately justify the request to purchase in excess of 8,000 dosage forms per month."³⁶⁶

9.5.6 Under the CSMP, threshold setting combined with other techniques resulted in a SOM program that continued to be non-compliant with the basic DEA requirements for controlled substances, as well as the terms of the company's 2008 settlement agreement.

McKesson replaced the LDMP program with the new Controlled Substance Monitoring Program or "CSMP" in 2008. The CSMP utilized a threshold system and for the first time created a mechanism by which orders could be blocked once that threshold was met in a given month. However, as the DEA repeatedly noted, the CSMP program consistently failed to highlight more than a few suspicious orders despite increasing levels of controlled substances purchases by McKesson customers.³⁶⁷ The DEA also expressed concern that McKesson in utilizing the CSMP failed to adhere to its obligation to "maintain a compliance program designed to detect and prevent diversion" as agreed in the 2008 settlement.³⁶⁸

[REDACTED]

³⁶³ See Email from Alexandra Feigel to Diane Martin, RE: November LDMP (Dec. 10, 2017) (Responding to D. Martin's email providing the November LDMP data), MCKMDL00540033.

³⁶⁴ *Id.*

³⁶⁵ *Id.*

³⁶⁶ Letter from J. Gilbert to L. Barber at 3 (Apr. 25, 2007) (Mr. Gilbert from Hyman, Phelps & McNamara was McKesson's outside counsel and Mr. Barber was Associate Chief Counsel in the Office of Chief Counsel, Diversion and Regulatory Litigation Section, DEA), MCKMDL00330924 (Also stamped MCK-HOI-002-0000001).

³⁶⁷ See Letter from D. Cutteman, *et al.*, to G. Hobart; Letter from W. Ihlenfeld, II 3/20/2014 letter to G. Hobart; J.F. Walsh 8/13/2014 letter to G. Hobart.

³⁶⁸ See 2008 MOA at 3, § II.1(a).

A. Establishing Thresholds

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

³⁶⁹ See CSMP Manual 2011 at 2, § 1.1.

³⁷⁰ *Id.*

³⁷¹ See CSMP Manual 2011 at 3, § 1.1.2

³⁷² See CSMP Manual 2013 at 2-3, § 1.1.

³⁷³ See Letter from J.F. Walsh to G. Hobart (Aug. 13, 2014) at 7; *see also* W. De Gutierrez-Mahoney Deposition at 249:11- 250:8.; Email from Tom McDonald to Tom Smith, *et al.*, RE: Clarifying the “partial” issue in CSMP (Jun. 10, 2010), MCKMDL00633917.

³⁷⁴ See ISMC CSMP Manual 2015 at 31, § 6.1.II.A.

³⁷⁵ *Id.* at 33.

³⁷⁶ *Id.* at 32-34 (Figure 7: AA00 Family Code -Default Thresholds).

³⁷⁷ See McKesson Regional Statistical Norms 2/24/2014 at 2 (By February 2014, the average number of doses (3 months rounded) was 24,500 units for both hydrocodone and oxycodone or more than 8,100 units per month); *see infra* LDMP discussion at section 3.2.5.2.

[REDACTED].³⁸⁵ The intent in setting thresholds this way was that most of McKesson's customers would never reach their established thresholds.³⁸⁶

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED].³⁸⁸

This justification does not make sense. [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED] In either case, McKesson was not in compliance with federal SOM requirements and expectations.

Furthermore, [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED] In August 2011, Dave Gustin, a Director of Regulatory Affairs ("DRA"), raised similar concerns about the thresholds being too high:

I have thought of an area that needs tightening up in CSMP, and it is the number of accounts we have that have large gaps between the amount of Oxy or Hydro they are allowed to buy (their threshold) and the amount they really need. (Their current purchases). The increases the 'opportunity' for diversion by exposing more product for introduction into the pipeline that may be being used for legitimate reasons.... If you look at the lists, you may be able to make the call to reduce some thresholds based on your or the sales person's knowledge of the acct."³⁹⁰

McKesson, however, appears not to have applied significant effort to address the threshold over-inflation until 2015, when this effort resulted in sizable threshold reductions for many customers.³⁹¹ Those sizable reductions

³⁸⁵ See W. De Gutierrez-Mahoney Deposition at 249:11 to 250:8; *see also* J.F. Walsh 8/13/2014 letter to G. Hobart at 7.

³⁸⁶ *See* T. McDonald 6/10/2010 email to Tom Smith, MCKMDL00633917.

³⁸⁷ Email from N. Hartle to M. Bishop, *et al.*, *Wakefern Threshold Methodology* (Sept. 9, 2014), MCKMDL00430124.

³⁸⁸ N. Hartle Deposition at 200:4-8.

³⁸⁹ Email M. Oriente to D. Walker, *et. al*, Regulatory Meeting 3/5 & 3/6, Attachment at 7 (Mar. 7, 2008), MCKMDL00545048 at MCKMDL00545054.

³⁹⁰ *See* Email from D. Gustin to D. Fagerskog, *et al.*, a couple of attachments were not sorted right. Here they are again. Project for the next few weeks., (Aug. 31, 2011), MCKMDL00507799.

³⁹¹ *See* Memorandum from N. Hartle to File, *McKesson's Controlled Substance Monitoring Program Oxycodone Threshold Reduction Report* (Feb. 9, 2015) (reducing oxycodone thresholds on 2,624 RNA customers), MCKMDL00402184; *see also* Email from N. Hartle

indicate that McKesson knew its thresholds were not set properly to identify suspicious orders and prevent potential diversion situations from occurring.

C. Threshold Disclosures

As part of the effort to ensure that the CSMP was customer-friendly and allowed “McKesson customer[s] [to] continue with business as usual,”³⁹² McKesson began preemptively telling customers they were approaching their threshold limits and at risk for order blocking.³⁹³ Doing so undercut the effectiveness of the thresholds as customers, whether diverting or not, were warned of the impending limit giving them time to request a threshold increase so that the threshold was not triggered. In fact, the following justification was stated internally for creating the threshold warning report in the first place:

We are in the business to sell the product. If we could produce a report ... that warned customers approach to the threshold, say at 85% of their 10,000 dosages, work could begin on justifying an increase in threshold prior to any lost sales.³⁹⁴

Helping customers avoid thresholds and order blocking are not objectives of a legitimate SOM program. Maintaining effective controls to prevent diversion, as well as identifying and reporting suspicious orders, should instead be the goal.

The standard for notification appears to be when the customer reached approximately 90% of their threshold.³⁹⁵ Although McKesson was not supposed to share the threshold with the customers, even if the customers were not told what their threshold was, they could back into the number with a bit of effort.³⁹⁶ The rationale for not telling customers exactly what their thresholds were was “so that they can’t try to manipulate a way around it, right, and get drugs from other suppliers or other distributors or something like that.”³⁹⁷

Customers, in fact, did not always have to back into their thresholds; they simply got them from their Distribution Center sales team. [REDACTED]

[REDACTED]

to K. Peck, FW: Threshold Reduction Initiative – 9143 and 9144 (May 1, 2005) (reductions for ISMC customers), MCKMDL00410744.

³⁹² Email attachment from D. Walker to Bill de Gutierrez-Mahoney, *McKesson Controlled Substances Monitoring Program – Program Guide for Pharmacies*, 2 (Apr. 4, 2008); *see also* Email D. Walker to #PGVDO, *et al.*, List 1 Chemicals (Jun 3, 2010) (“There will no longer be “partial” omits of controlled substances or List 1 chemicals. If a customer exceeds their threshold on a certain item the entire item ordered will not be shipped.”), MCKMDL00633917 at MCKMDL0063919.

³⁹³ *See* CSMP Manual 2011 at § 2.

³⁹⁴ *See* Email from S. Mackarness to G. Hillard (Oct. 26, 2006), MCKMDL00543971 at MCKMDL00543972.

³⁹⁵ *See* N. Hartle Deposition at 135:7-13.

³⁹⁶ *See* N. Hartle Deposition at 136:1-8.

³⁹⁷ *See* N. Hartle Deposition at 134:15-18.

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

For example in April 2013, Amanda Miller, RNA Account Manager in Support Solutions, sent Melanie Petropoulos, Vice President, Pharmacy for Marc Glassman, Inc. a copy of the internal system print out showing their oxycodone threshold, the monthly purchases on record and the percentage of thresholds used for three Glassman entities.⁴⁰¹ This was not an isolated instance,⁴⁰² as the sharing of thresholds started almost as soon as the CSMP program was instituted.⁴⁰³

It also seems that Regulatory Affairs did not always follow the rule that thresholds should not be shared. For example, in April 2008, Mr. Walker, head of the SOM program, wrote in an email that he was not opposed to honor a customer's request for "a breakdown of our current thresholds on our controlled substances for each site" because he was sure that "we will get a lot of these requests from customers wanting to make sure we have it covered."⁴⁰⁴

Overall, McKesson's "proactive" approach of communicating threshold warnings to customers effectively rendered the thresholds meaningless as a diversion prevention goal. This was further exacerbated by the policy of allowing Distribution Center personnel to contact and discuss the actual threshold amounts with customers. Thus, the employees the CSMP program who were counted on to be the "eyes and ears" of the program and who were on the front lines, were rewarded not for spotting and stopping suspicious orders, but for making sure the SOM program did not stop customer orders and disrupt business, thereby undermining the CSMP as a diversion prevention control.

³⁹⁸ See CSMP Manual 2011 at § 2.1.

³⁹⁹ See *id.*

⁴⁰⁰ N Hartle Deposition at 138:1-16.

⁴⁰¹ See Email from Amanda Miller to Melanie Petropoulos, CSMP 4/23/13 (Apr. 23, 2013), MCKMDL00476776.

⁴⁰² See, e.g., Email from Denise Joslyn to Joseph Lahvoich, Acme 1/11/2013 CSMP (Jan. 11, 2013), MCKMDL00496271; Ned McKenna, *et al.*, *CVS CSMP: Threshold Review*, 9 (Jan. 9, 2009), MCKMDL00574488 at MCKMDL00574496 ("McKesson will manage reasonable threshold requests for CVS as needed..."); Email from Sabrina Cook to Gregory Carlson, Giant Eagle CSMP Thresholds (Oct. 22, 2008), MCKMDL00628660.

⁴⁰³ See Sabrina Cook 10/22/2008 email to Gregory Carlson.

⁴⁰⁴ Email from D. Walker to W. De Gutierrez-Mahoney, *et al.*, FW: NEW DEA ORDERING STANDARDS (Apr. 17, 2008), MCKMDL00543610.

D. Exceeding the Threshold (TCRs and Level 1-3 Review)

Similarly, McKesson's program and practices also undercut the controls associated with when orders are deemed "suspicious" (e.g., order blocking). [REDACTED]

[REDACTED]
[REDACTED]
[REDACTED] Exceeding the threshold also triggered the SOM Level 1-3 review process, which was adopted from the LDMP.⁴⁰⁷

Although conceived as a controlled, documented way to make appropriate adjustments to thresholds, the Threshold Change Request ("TCR"), like any internal control, could be abused, which it was in McKesson's case. For any control, once the exceptions or changes become commonplace, the control is rendered ineffective. In the case of thresholds, TCRs were commonplace at McKesson.⁴⁰⁸ The solution when this occurs is to change the control or disallow exceptions, not to simply keep processing exception requests.

[REDACTED]
[REDACTED]
[REDACTED]

[REDACTED]
[REDACTED]
[REDACTED]

[REDACTED]
[REDACTED] Therefore, it was a relatively common practice for Distribution Center personnel to contact the customer upon receiving a customer threshold warning and put through a TCR.⁴¹⁴ For example, as discussed above, Amanda Walker provided the VP of Pharmacy for Marc Glassman, Inc. their threshold numbers and current purchasing levels with the brief note "Let me know if you need to

⁴⁰⁵ See CSMP Manual 2011 at § 2.2; see also D. Walker 6/3/2010 email to #PGVDO, *et al.* ("There will no longer be "partial" omits of controlled substances or List 1 chemicals. If a customer exceeds their threshold on a certain item the entire item ordered will not be shipped.").

⁴⁰⁶ *Id.*

⁴⁰⁷ See CSMP Manual 2011 at §§ 2.2.1-2.2.4; N. Hartle Deposition at 140:4-16.

⁴⁰⁸ See Email A. Miller to D. Graziano, RE: CSMP 4/23/13 (Apr. 24, 2013), MCKMDL00476776.

⁴⁰⁹ See CSMP Manual 2011 at § 1.3.1.

⁴¹⁰ *Id.* at § 1.3.1

⁴¹¹ See CSMP Manual 2011 at § 1.3.2.

⁴¹² See Email from A. Miller 4/24/2018 to D. Graziano.

⁴¹³ N Hartle Deposition at 125:9-24.

⁴¹⁴ See S. Mackarness 10/26/2006 email to G. Hillard ("If we could produce a report ... that warned a customers [sic.] approach to the threshold ... work could begin on justifying an increase in threshold prior to any lost sales.").

make any adjustments.”⁴¹⁵ As was the case with threshold disclosures, using TCRs to avoid triggering the suspicious order process through continually raising the bar was widespread.⁴¹⁶

By using Distribution Center personnel to solicit in advance whether a threshold increase was needed and to facilitate TCRs, McKesson eviscerated its own diversion control by ensuring that customers avoided triggering order blocking, as well as suspicious order reporting and investigations. In short, McKesson employees used TCRs to manipulate the SOM process to prevent lost sales.⁴¹⁷

E. Customer Due Diligence or Level 1 Review

.⁴¹⁹ However, although McKesson's due diligence process looked functional on paper, McKesson's implementation of the program made it a *pro forma* exercise.

McKesson's interactions with Tug Valley Pharmacy aptly illustrate the problems with this approach. In May 2015, McKesson received a new customer questionnaire from Tug Valley in which the pharmacy disclosed that Miami-Luken had terminated its ability to purchase controlled substances.⁴²³ McKesson also learned that Tug Valley was involved in pending litigation relating to allegations of controlled substances diversion.⁴²⁴ However, despite this knowledge, McKesson approved Tug Valley's application in July 2015.⁴²⁵ In January

⁴¹⁵ See Amanda Miller 4/23/2013 email to Melanie Petropoulos.

⁴¹⁶ See J.F. Walsh 8/13/2014 letter to G. Hobart at 12.

⁴¹⁷ See S. Mackarness 10/26/2006 email to G. Hillard.

⁴¹⁸ See, e.g., James Arnold 6/2013 presentation at slides 42-53); Diane Murphy, *The Federal Sentencing Guidelines for Organizations: A Decade of Promoting Compliance and Ethics*, 87 Iowa L. Rev. 697, 703 (2002).

⁴¹⁹ See CSMP Manual 2013 and CSMP Manual 2011 §§ 1.3.1, 2.2 and 3.

⁴²⁰ CSMP Manual 2013 at § 1.2.2.

⁴²¹ See CSMP Manual 2013 at § 3.

⁴²² CSMP Manual 2013, at, § 3.2.1 and § 3.2.2.

⁴²³ See W.Va. Red Flags Report at 142.

⁴²⁴ *Id.* at 147

425 *Id.*

2016, McKesson suspended Tug Valley after a *CBS News* report focused on the role distributors in the opioid crisis and “prominently featured Tug Valley”.⁴²⁶ Tug Valley submitted a new application in February 2016 representing that the pharmacy was under new ownership, and while the questionnaire lacked answers to many of the crucial questions, the McKesson Regulatory Investigative Report found no “red flags.”⁴²⁷ McKesson also learned that the former owner still had a security interest in the pharmacy, which supposedly was under new ownership.⁴²⁸ Once more, despite the existence of new “red flags,” McKesson reinstated the pharmacy.⁴²⁹

The failure to conduct meaningful oversight or challenge “flimsy rationales” was systemic across the McKesson network as confirmed on multiple occasions by DEA and DOJ officials in various communications.⁴³³ McKesson, itself recognized the issue, but allowed these practices to continue without ensuring appropriate changes occurred with predictable results; decisions to increase customer thresholds were made without an apparent rational basis.⁴³⁴

⁴²⁶ *Id.* at 145 and 147.

⁴²⁸ *Id.* at 153.

⁴³⁰ See ISMC CSMP Manual 2015 at 18-22.

⁴³² See DEA 9/27/2006 Letter at 3.

⁴³⁴ See, e.g., Email from D. Gustin to D. Fagerskog, *et al.*, CSMP Contribution, DCM Call, Tightening up our increase process (Apr. 15, 2011) (“We also need to tighten up the process regarding granting increases. We have gotten to the point where certain % of increase [sic.] are almost automatic and where we are too easily accepting of ‘reasons’ like ‘business increase’ for raising thresholds by small amounts.”), MCKMDL00507221 at MCKMDL00507223; *see also* B. Russell email to B. de Gutierrez-Mahoney, *et al.*, *Threshold Change Requests* (Jul. 27, 2012) (citing email from T. McDonald in Regulatory Affairs on how to appropriately document TCRs); MCKMDL00633455.

This persistent lack of oversight of customer behavior perhaps is best illustrated by the New Castle, Pennsylvania distribution center, which served parts of West Virginia as well as Cuyahoga and Summit Counties in Ohio. From 2000 to the present, Blaine Snider, a 39-year McKesson employee, has been New Castle's Director of Operations.⁴³⁵ Mr. Snider, as the Director of Operations or Distribution Center Manager ("DCM"),⁴³⁶ was supposed to raise concerns about potentially suspicious customer orders and ultimately could refuse shipments to those customers he believed were placing unjustifiable suspicious orders.⁴³⁷ He has, in fact, done neither during his tenure.

1. The "Thanksgiving" Increases

Perhaps the most egregious example of failed oversight and due diligence occurred in December 2008 and involved Mr. Snider, Dave Gustin, DRA and Michael Bishop. On November 28, 2008, a request was sent to change more than 200 RNA account thresholds to Micheal Bishop, Compliance Analyst Business Process, RNA Support Solutions.⁴³⁸ The form lists "various RNA customers" including Wal-Mart, Rite Aid and Target on an attachment and lists the controlled substances requested as "various."⁴³⁹ A permanent 30% threshold increase was requested and the reason for the change was listed as, "Increase due to Thanksgiving holiday – 30% increase." When asked about the Thanksgiving holiday increases in his deposition, Mr. Snider testified it was McKesson's policy to grant permanent threshold increases based on holidays.⁴⁴⁰

Mr. Bishop, in turn, passed the request to Dave Gustin, Director of Regulatory Affairs, who simply made the changes before processing the paperwork. As a result, Mr. Gustin emailed Bishop on December 16, 2008, requesting that Bishop provide a "TCR from you signed and dated the 30th. I will use it for the 30% increases I made for the RNAs that day after you emailed me all those reports."⁴⁴¹ On December 17, 2008, Mr. Gustin confirmed with Bishop that these were the "Thanksgiving increases" that occurred on November 28th.⁴⁴²

Not only was this 30% increase put through as a permanent versus perhaps a temporary request, but also the rationale of the Thanksgiving holiday was arbitrarily applied to 200 accounts by the DRA. To justify his actions in not performing the required reviews prior to making the changes, Mr. Gustin wrote that he "was the only DRA on and so my time was spent making the changes."⁴⁴³ Furthermore, it appears that his request to "backdate" the TCR form as part of a larger and more commonplace practice within McKesson's Regulatory Affairs function. According to Nathan Hartle, "it's standard to document things after. You may make the

⁴³⁵ See B. Snider Deposition at 19:4-13. According to Snider, the terms Director of Operations and Distribution Center Manager ("DCM") are used interchangeably at McKesson.

⁴³⁶ See *id.* at 121:1-4.

⁴³⁷ *Id.* 108:2-17 and 109:17:22.

⁴³⁸ See Blaine Snider, *Threshold Change Request Form*, (Nov. 28, 2008), MCKMDL00363951 at MCKMDL00363954.

⁴³⁹ See *id.*; Attachment is found at MCKMDL0036953.

⁴⁴⁰ See B. Snider Deposition at 223:22-24 and 224:1-10.

⁴⁴¹ See Dave Gustin email to Michael Bishop, RE: could you do me a favor (Dec. 16, 2008), MCKMDL00000522.

⁴⁴² See Dave Gustin email to Michael Bishop, RE: could you do me a favor (Dec. 17, 2008), MCKMDL00000521.

⁴⁴³ See Dave Gustin email to #PGDCM, *et al.*, FW: could you do me a favor (Dec. 17, 2008), MCKMDL00363951 at MCKMDL00363955.

decision based on the information you have in your own notes. To put into the format may happen after that. That's not uncommon.”⁴⁴⁴ Mr. Hartle also stated, “you make decisions based on your notes and do the official documentation later.”⁴⁴⁵ To be clear, “backdating” is never acceptable, especially when it involves regulatory documentation. Furthermore, the “Thanksgiving” threshold increases were not the only instance where improper due diligence was conducted concerning controlled substance orders by McKesson customers.

2. Retail National Account Deference

The Thanksgiving increases are symptomatic of a broader pattern of conduct by McKesson that involved deferring to the retail national account (“RNA”) customers to police their own pharmacies, thereby mistakenly absolving McKesson of the need to apply its SOM program to those accounts. For example, within the documents produced by McKesson, there are numerous examples of Retail National Account (“RNA”) stores in Summit and Cuyahoga Counties being granted threshold increases with little or no investigation or interrogation.⁴⁴⁶

McKesson’s actions with RNA customers run contrary to its regulatory obligations and are inconsistent with the actions of a reasonable and prudent distributor because controlled substances compliance is not deferrable or delegable.

McKesson’s Senior Director of Distribution Operations, Donald Walker, readily acknowledged that McKesson did not ask for dispensing data in order to verify the legitimacy of threshold increases for retail national account customers and generally deferred to those customers to decide when it was appropriate for them to get threshold increases for controlled substances.⁴⁴⁷ For example, as seen in a January 2009 presentation, McKesson outlined its plan for automatic threshold increases for CVS stores when they approached their threshold and to only seek a justification for thresholds increases from CVS if the increases were “extraordinary” and without “further CVS explanation.”⁴⁴⁸ McKesson’s erroneous reasoning for such automatic threshold increases was to “minimize disruption of business,” and to ignore reviewing “routine” threshold increases.⁴⁴⁹

9.5.7 McKesson’s Internal Audit process was ineffectual and not a reasonable control to detect non-compliance.

McKesson’s Board of Directors in April 2018 publicly released a report by a Special Committee of the Board that examined the company’s oversight of opioid distribution by senior management and the Board during the

⁴⁴⁴ See N. Hartle Deposition at 166:1-6

⁴⁴⁵ See N. Hartle Deposition at 165: 9-11.

⁴⁴⁶ See e.g., Giant Eagle 4030 Due Diligence File, MCKMDL00555448; Email from S. Cook to D. Gustin, *et al.*, RE: Pain mgt, (Nov. 1, 2010), MCKMDL00512974; Email from D. Gustin to S. Cook, FW: Giant Eagle CSMP Thresholds, (Feb. 23, 2009), MCKMDL00628614; Giant Eagle 6376 Due Diligence File MCKMDL00555473; Giant Eagle 0209 Due Diligence File MCKMDL00555484.

⁴⁴⁷ See Donald Walker Deposition, 190-193 (Jan. 10, 2019).

⁴⁴⁸ See Presentation by N. McKenna, *et al.*, CVS CSMP: Threshold Review, 2 (Jan. 9, 2009), MCKMDL00574488.

⁴⁴⁹ *Id.*

time period between the two settlements with the federal government (2008 to 2017).⁴⁵⁰ The Special Committee concluded that:

the Committee's investigation revealed a strong moral culture at McKesson, as led by Senior Management and reinforced by the Board, and that the Company's Chief Executive Officer and others in Senior Management created a strong tone at the top of McKesson that encouraged ethical and compliant conduct.⁴⁵¹

The Special Committee felt that the Audit Committee's and Senior Management's belief that the compliance program was satisfactory and working effectively was reasonable because the company "had oversight procedures in place, including Internal Audit reviews of the Company's compliance program and distribution facilities" and that there were "positive Internal Audit review results (including findings that management had addressed, or was addressing, any identified issues)."⁴⁵² My review of the details, however, revealed a very different picture.

I reviewed actual Corporate Internal Audit ("IA") reports of the controlled substances program and below provide a detailed discussion on those that occurred in 2007, 2010 and 2012.⁴⁵³ As the Special Committee noted, for those audits the overall audit ratings were Green-Satisfactory, Yellow-Need Improvement, and Green-Satisfactory respectively.⁴⁵⁴ However, in all three reports, IA either failed to probe the area deeply enough or failed to understand the importance of what they observed. In all three cases the risks were downplayed, which led to an overall audit rating and Executive Summary that conveyed a false sense that McKesson's SOM program was in compliance.

A. 2007 Report

In the 2007 report, the IA team noted issues with the LDMP program (Observation #1), the DEA Licensure Process (Observation #2) and Licensing Requirements policies and procedures (Observation #6).⁴⁵⁵ All three observations involved a lack of formal processes, or guidance or training resulting in McKesson employees not knowing what was required of them.⁴⁵⁶ All three observations were classified as being of "moderate" significance and none made it into the executive summary. This is in spite of the fact that a basic requirement of a controlled substances compliance program is to ensure that customers are lawfully entitled to receive shipments of controlled substances and that exceeding thresholds rendered orders potentially "suspicious" and

⁴⁵⁰ See MCK Teamsters Response at 4.

⁴⁵¹ *Id.* at 26.

⁴⁵² *Id.* at 28 and 29-30.

⁴⁵³ See J. Robinson, *Audit Report- DEA Licensure and LDMP Audit U.S. Pharmaceuticals*, 08-SSPH-04 (Oct. 9, 2007), MCKMDL00591251; M. Fuller, *Audit Report – Distribution Center and Controlled Substances Monitoring Program U.S. Pharmaceutical Distribution Operations*, 10-SSPH-11 (Jul. 20, 2010), MCKMDL00591972; M. Fuller, *Audit Report – Distribution Center Audit U.S. Pharmaceutical Distribution Operations*, 12-SSPH-08 (Apr. 10, 2012), MCKMDL00594099. ["IA Report (year)"].

⁴⁵⁴ *Id.*

⁴⁵⁵ See IA Report 2007 at 7-9.

⁴⁵⁶ *Id.*

thus reportable to the DEA.⁴⁵⁷ In other words, being out of compliance with DEA regulations or not knowing them at all was not deemed a critical observation worthy of senior management's or the Board's attention.

B. 2010 Report

The 2010 report highlighted the fact that four Distribution Centers (Memphis, Lakeland, Phoenix, and Landover) were not following required processes in that required TCRs and Omit Reports were not being signed on a regular basis by the DCMs, which as the auditors noted, called into question whether the reviews were even being done.⁴⁵⁸ However, just like the 2007 report, the observation was deemed to be of "moderate" significance and was not highlighted in the executive summary even though threshold usage and monitoring were parts of the 2008 DEA settlement. The action plan from each Distribution Center was essentially "we'll do better" and make sure the paperwork is signed, which is a wholly inadequate response.⁴⁵⁹

Finally, it appears that IA confined its review to simply looking to see if the documents were signed and dated as required. There is no evidence that any of the rationales were examined to see if they passed a commonsense test (i.e., no Thanksgiving holiday rationales for permanent increases). Therefore, IA apparently undertook a limited transactional review.

C. 2012 Report

[REDACTED]

Once more the observation earned a "moderate" significance score and was not highlighted in the Executive Summary. This occurred even though the observation and the Distribution Center responses were the same as in 2010 but involved different distribution centers. This fact should have triggered heightened concern that the documentation concerns were systemic and not being remedied by U.S. Pharma Regulatory Affairs. Once more it appears that IA did not examine any underlying rationales for the TCRs or the adequacy of the Level 1 data.

The deficiencies in the IA reports are such that it was readily apparent why the IA ratings of the controlled substances compliance program are diametrically opposed to what the DEA repeatedly uncovered and brought to McKesson's attention in the same time period. Thus, the IA process was ineffective and failed to provide senior management and the Board with a true picture of how the controlled substances compliance program was operating.

⁴⁵⁷ See 21 C.F.R. §1301.74(a) and (b).

⁴⁵⁸ See IA Report 2010 at 9-10 (Observation #4).

⁴⁵⁹ *Id.*

⁴⁶⁰ See IA Report 2012 at 9-10 (Observation #4).

⁴⁶¹ *Id.* at 9.

9.5.8 McKesson failed to undertake appropriate corrective actions to ensure its controlled substances program was compliant with its regulatory obligations.

Although the DEA repeatedly placed McKesson on notice before the 2008 settlement of its failures to maintain a SOM system, McKesson failed to rectify the situation. Nor did the company correct the situation after the 2008 settlement, which required the DEA to undertake a second enforcement action in 2017. This type of recidivist behavior and reoccurrence of similar misconduct, as noted by the FSGs, “creates doubt regarding whether the organization took reasonable steps to meet the requirements of this guideline” or in this case of the requirements of the CSA and its accompanying regulations.⁴⁶² It certainly makes clear that McKesson lacked an adequate corrective action process.

The same lack of an appropriate corrective action process is illustrated by the Internal Audit reports. Given the fact that IA continued to perform the same type of audits in the same way in the face of DEA’s diametrically opposite findings, indicates that McKesson made little or no attempt to ascertain why IA and DEA were getting such different results and to make any necessary course corrections.

However, what is more troubling is that McKesson, beyond rewriting the CSMP manual, ultimately increased the program’s complexity by splitting it in half and by only making other relatively minor alterations. In short, it seems McKesson did not undertake the more systemic actions needed to eradicate its previous culture. The Special Committee’s 2018 report amply supports this lack of any “decisive steps” stating “the Committee’s investigation revealed a strong moral culture at McKesson, as led by Senior Management and reinforced by the Board, and that the Company’s Chief Executive Officer and others in Senior Management created a strong tone at the top of McKesson that encouraged ethical and compliant conduct.”⁴⁶³ The evidence reviewed in this report does not support that conclusion.

The Board continues to maintain that McKesson’s senior management “attempted in earnest to meet the DEA’s suspicious order reporting regulations, despite a lack of specific instructions or feedback as to the parameters of a program that the DEA would find acceptable.”⁴⁶⁴ Between the CSA, the DEA regulations, the general industry letters, the numerous private and specific letters to the company, and the 2008 Settlement Agreement, the DEA was clear about what its concerns were, what the Agency felt were McKesson’s responsibilities, and where it felt McKesson was lacking. McKesson, however, did not decisively rectify matters. Therefore, with the 2017 settlement, DEA essentially removed McKesson’s discretionary ability to design and operate the controlled substances program and tailor it to fit McKesson’s company structure. Instead, the DEA substituted a highly detailed government-designed roadmap set out in the Compliance Addendum.

Therefore, I believe the answers to the questions outlined by McKesson’s Board of Directors in terms of McKesson’s senior management should be:⁴⁶⁵

⁴⁶² See FSGs 2004 at § 8B2.1, Application Note 2(D).

⁴⁶³ See MCK Teamsters Response at 26.

⁴⁶⁴ See *id.* at 28.

⁴⁶⁵ See MCK Teamsters Response at 5.

- **Did McKesson's senior management intentionally disregard, or turn a blind eye to, the Company's compliance obligations following the 2008 settlement?** There is enough evidence to demonstrate that senior management "turned a blind eye" by not providing adequate support, commitment, resources, or accountability regarding its opioid distribution operations, despite its 2008 agreement with the DEA and despite being put on notice to those deficiencies by the DEA between 2008 and 2017.
- **Did senior management act in bad faith or recklessly in connection with the Company's distribution of opioids?** At a minimum, senior management failed in its duty to exercise appropriate oversight and governance over the company's distribution of opioids. These failures were systemic and not confined to a few "rogue" Distribution Centers or "bad apple" employees, but instead permeated the entire program.
- **Did McKesson's senior management take advantage of the opioid crisis by encouraging or condoning shipments of opioids to customers it knew or should have known were diverting the drugs for illegitimate use?** Senior management clearly knew or should have known that many McKesson customers were purchasing opioids at ever increasing levels for which no rational basis for the increases had been established. Furthermore, when confronted by the DEA with evidence that this was happening, senior management persisted with a program it knew or should have known was defective and out of compliance with its regulatory obligations. In short, the company simply never looked for legitimate explanations the ordering patterns it encountered.
- **Did the Company prioritize revenue over compliance?** In my opinion, McKesson's continuing tacit endorsement of minimal compliance prioritized allowing its customers to continue "doing business as usual" (e.g., selling more opioids) over meeting its regulatory and compliance oversight responsibilities. As one McKesson Vice President simply noted, "[w]e are in the business to sell product."⁴⁶⁶ Such widespread failures, including not equating good compliance with good business, in my experience, can only happen because of this type of an inappropriate "tone at the top."

Finally, as noted in the Internal Audit discussion, the Board's Special Committee and its outside law firm, even with the benefit of hindsight knowing what the DEA found, failed to identify and address the problem that important observations about the controlled substances compliance program in IA reports were not being highlighted for the Board. Therefore, despite an IA process which provided an inaccurate view of the state of the SOM program, the Board continued to see the IA process as a significant factor justifying its basic conclusion of no wrongdoing by McKesson senior officials, including Mr. Walker, the SVP in charge of the program from 1997 to 2015.

9.5.9 No evidence was presented that McKesson had a formal risk assessment process during the period.

Without belaboring the point, McKesson also did not implement an adequate or formal risk assessment process involving controlled substances distribution. In fact, no evidence was uncovered in this review to indicate that

⁴⁶⁶ See Email Sharon Harkness to Gary Hillard, *RE: hydrocodone reports* (Oct. 26, 2006), MCKMDL00543971 at MCKMDL00543972.

McKesson ever established a formal risk assessment process during the review period. Had there been such a process, McKesson would have made changes and improvements to its controlled substances program on its own. However, as can be seen with the LDMP and the various iterations of the CSMP, McKesson only made changes in response to DEA's threat of initiating actual enforcement actions. Had there been such a proactive risk assessment process, the IA findings in the 2007, 2010, and 2012 reports would not have been deemed to be of "moderate" significance.

9.6 Program Changes (2014 to Present)

9.6.1 The AGI Engagement

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

- [REDACTED]
- [REDACTED]
- [REDACTED]

- [REDACTED]

- [REDACTED]
 - [REDACTED]

[REDACTED]

[REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]

⁴⁶⁷ See MCK Teamsters Response at 24.

⁴⁶⁸ See Analysis Group, Inc., *Suspicious Order Monitoring Threshold System for McKesson Independent Retail Customers – Description and Rationale*, 2 (May 12, 2017), MCKMDL00437057 [“AGI SOM Description”].

469 *Id.*

⁴⁷⁰ *Id.* at 5-6.

⁴⁷¹ See AGI SOM Description at 2.

⁴⁷² *Id.* at 3.

[REDACTED]

[REDACTED]
[REDACTED]
[REDACTED]
[REDACTED]
[REDACTED]
[REDACTED]
[REDACTED]

[REDACTED]
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[REDACTED]

[REDACTED]
[REDACTED]
[REDACTED]
[REDACTED]

[REDACTED]
[REDACTED]
[REDACTED]

⁴⁷³ *Id.* (emphasis in the original).

⁴⁷⁴ *Id.* at 4.

⁴⁷⁵ *Id.*

⁴⁷⁶ *Id.* at 5.

⁴⁷⁷ *See* AGI SOM Description at 8 and 10.

⁴⁷⁸ *See* AGI SOM Description at 7.

⁴⁷⁹ *Id.* (emphasis added).

481

[REDACTED]

480 *Id.*

⁴⁸² See Presentation by Krista Peck, *US Pharma CSMP Update*, 5 (Dec. 1, 2015), MCKMDL00459350.

⁴⁸⁴ See *id.* at 7.

⁴⁸⁶ See Presentation by Nate Hartle & The Analytics Team, Introduction [to] the ISMC Solver in Tableau, (Oct. 24-26, 2016), MCKMDL00427408.

⁴⁸⁸ McKesson Corporation's Third Supplemental Objections and Responses to Plaintiffs' Combine Discovery Requests at p. 9.

9.7 Accountability - Consistent Enforcement

McKesson, throughout the Review Period, failed to maintain and utilize appropriate disciplinary mechanisms either towards:

- customers, who routinely exceeded established SOM thresholds, or internal employees, who collaborated with customers to circumvent the threshold controls (i.e., “gamed the system”), or
- their supervisors, who either encouraged or ignored the inappropriate collaboration between McKesson employees and customers and
- their supervisors who failed to make the necessary improvements to the controlled substances compliance program for the program to meet McKesson’s regulatory obligations as a controlled substances distributor.

McKesson also did not use reasonable efforts to avoid putting “bad actors” in substantial authority positions over its distribution of controlled substances. Thus, there simply was no evidence presented of there being real consequences for circumventing or ignoring SOM controls.

9.7.1 Despite repeated breaches of company policies and DEA SOM requirements, McKesson failed to discipline those involved.

McKesson's overall compliance efforts (controlled substances and corporate compliance) also were ineffective because the company failed to hold those involved with serious breaches of the federal controlled substances requirements accountable, and in at least one case even promoted one of the culpable individuals to a position of greater authority within the controlled substances program. [REDACTED]

⁴⁸⁹ See W. Ihlenfeld letter to G. Hobart, *McKesson Matter – Ongoing Settlement Discussions*, (Mar. 20, 2014).

9.7.2 McKesson also failed to consistently terminate customers found in repeated breach of the SOM requirements.

While there are numerous examples of this occurring, some of which are discussed earlier in this report, a case in point is McKesson's sales to [REDACTED] as documented by the U.S. Attorney's Office for the Northern District of West Virginia. According to the U.S. Attorney's Office, McKesson in April 2012 notified the DEA that it had identified [REDACTED] as a suspicious customer and ceased sales of oxycodone to the pharmacy.⁴⁹⁶ However, according to the government, [REDACTED]

Cardinal Health, Inc.

⁴⁹¹ See *id.* at 25:3-10.

⁴⁹² See *id.* at 34:14-19.

⁴⁹³ See 2008 MOA.

⁴⁹⁴ See D. Cutteman 11/4/2014 letter to G. Hobart at 4.

⁴⁹⁶ See W. Ihlenfeld letter to G. Hobart, *McKesson Matter – Ongoing Settlement Discussions*, (Mar. 20, 2014) at 4.

⁴⁹⁷ *Id.* at 5.

⁴⁹⁸ *Id.* at 5.

10 Cardinal Health, Inc.

10.1 Background

Formed in 1979, Cardinal Health, Inc. (“Cardinal” or “Cardinal Health”) describes itself as a “global, integrated healthcare services and products company providing customized solutions for hospitals, healthcare systems, pharmacies, ambulatory surgery centers, clinical laboratories and physician offices.”⁴⁹⁹ Cardinal is headquartered in Dublin, Ohio and employs “nearly 50,000 people” and has annual revenues of \$137 billion.⁵⁰⁰

From a controlled substances perspective, Cardinal serves more than 26,000 pharmacies, and in the case of West Virginia, it was the largest supplier of controlled substances from 2005 to 2016.⁵⁰¹ As of its 2008 settlement with the government, Cardinal operated 27 distribution centers that handled controlled substances.⁵⁰² By 2012 that number had increased to 28.⁵⁰³

Cardinal Health’s controlled substance compliance program also has changed over time correlating to various regulatory enforcement milestones. Beginning in the fourth quarter 2007 and continuing into the first quarter of 2008, Cardinal received four Orders to Show Cause and Immediate Suspension of Registration (“ISOs”) from the DEA that ultimately resulted in the 2008 AMOA.⁵⁰⁴ In 2012, Cardinal Health again entered into a settlement agreement with the government.⁵⁰⁵ As discussed in detail below, at each enforcement milestone, Cardinal modified its program in response to those proceedings.

10.2 Executive Summary

My review of Cardinal’s controlled substances compliance efforts from 1996 to 2008 reveals a consistent pattern of systemic failure to meet its regulatory and corporate governance obligations with respect to controlled substances. After the 2008 government settlement Cardinal expended energy in generating a controlled

⁴⁹⁹ See CARDINAL HEALTH, Fiscal 2018 Form 10-K, 3 (Sept. 2018).

⁵⁰⁰ See CARDINAL HEALTH FACT SHEET, *Essential to care* (2018).

⁵⁰¹ See U.S. House Energy & Commerce Committee Report, *Red Flags and Warning Signs Ignored: Opioid Distribution and Enforcement Concerns in West Virginia*, 115th Cong., 243 (Dec. 19, 2018) [“W.Va. Red Flags Report”].

⁵⁰² See Administrative Memorandum of Agreement between the U.S. Department of Justice, Drug Enforcement Administration and Cardinal Health, Inc., 1 (Oct. 2, 2008), CAH_MDL_PRIORPROD_DEA12_00014414 [“2008 CAH AMOA”].

⁵⁰³ See Administrative Memorandum of Agreement between the U.S. Department of Justice, Drug Enforcement Administration and Cardinal Health, Inc., 1 (May 14, 2012), CAH_MDL2804_02465982 [“2012 CAH AMOA”].

⁵⁰⁴ See Letter from M. Leonhart, Order to Show Cause and Immediate Suspension of Registration, Auburn, Washington Distribution Center (Nov. 28, 2007), CAH_MDL_PRIORPROD_DEA12_00014414 at CAH_MDL_PRIORPROD_DEA12_00014430; Letter from M. Leonhart, Order to Show Cause and Immediate Suspension of Registration, Lakeland, Florida Distribution Center (Dec. 5, 2007), CAH_MDL_PRIORPROD_DEA12_00014414 at CAH_MDL_PRIORPROD_DEA12_00014434; Letter from M. Leonhart, Order to Show Cause and Immediate Suspension of Registration, Swedesboro, New Jersey Distribution Center (Dec. 7, 2007), CAH_MDL_PRIORPROD_DEA12_00014414 at CAH_MDL_PRIORPROD_DEA12_00014439; Letter from J. Rannazzisi, Order to Show Cause Stafford, Texas Distribution Center (Jan. 30, 2008), CAH_MDL_PRIORPROD_DEA12_00014414 at CAH_MDL_PRIORPROD_DEA12_00014444, [“[Facility Name][Year]ISO” or collectively “2007 ISOs”].

⁵⁰⁵ See 2012 CAH AMOA.

substances compliance program. However, Cardinal's poor program design exacerbated by poor implementation ultimately defeated those efforts resulting in another government settlement agreement in 2012. Thus, Cardinal created a "paper program" that was neither effective for controlling diversion or identifying and reporting suspicious orders of controlled substances.

The program's failure cannot be attributed to one root cause. Both corporate culture and Cardinal's overreliance on technology played prominent roles. Cardinal's controlled substances compliance efforts did not have the necessary support and commitment of senior management and its Board of Directors. Despite the well-documented existence of the applicable compliance program standards for controlled substances and corporate compliance dating back to the 1970s; Cardinal did not make a concentrated effort to comply with those until 2008. Nor did Cardinal adopt a proactive approach towards compliance, even after its settlement in 2008. Cardinal's management simply did not act unless it was pushed to do so by the DEA, and then the company only did the minimum.

In the case of technology, Cardinal placed a premium on its analytical systems to detect suspicious orders and potential diversion, while neglecting the importance of the human element in making sense from the data outputs.

Despite creating the appearance of a documented and adequate suspicious order monitoring program, the reality was the SOM and anti-diversion programs simply did not work. Even George Barrett, Executive Chairman of the Board for Cardinal Health, reluctantly admitted as much. When asked by a House Energy and Commerce subcommittee whether Cardinal inquired about the reason for the higher drug order when a pharmacy goes over its monthly drug threshold, Barrett replied:

"I think the thresholds probably should have been set with a different set of eyes. I've mentioned this notion of asking different questions. And I think today we'd probably set those quite differently. But I think at the time of those pharmacies you referred to, thresholds probably should have been adjusted down more quickly."⁵⁰⁶

When pressed further about whether Cardinal simply turned a blind eye to the impact of requested threshold increases even when the rationale did not make sense, Barrett responded:

"I have no reason to question the good intent of those doing that kind of assessment. They were professionals. I think they were looking at the incoming order of prescribing. I think now we know some of that prescribing was driven by some behavior that we would have liked to have caught in the physician world."⁵⁰⁷

The program failed in part because Cardinal failed to embrace its affirmative duties under the CSA to create and maintain an effective program to prevent diversion. It also failed because Cardinal developed a program that was so convoluted and riddled with loopholes that Cardinal could avoid identifying orders as suspicious and continued supplying customers that it knew or should have known were engaging in diversionary behavior.

⁵⁰⁶ W.Va. Red Flags Report at 223, *citing* Combating the Opioid Epidemic: Examining Concerns About Distribution and Diversion: Hearing Before the Subcomm. on Oversight and Investigations of the H. Comm. on Energy and Commerce, 115th Cong at 108 (2018).

⁵⁰⁷ *Id.*

10.3 Impact

Cardinal's activities in West Virginia are indicative of the results caused by these compliance failures. In the case of West Virginia, Cardinal told the House Energy and Commerce Committee ("E&C Committee") that it could not determine if the company submitted any suspicious orders from 2006 to 2012 to the DEA involving customers in the state, but from 2012 to 2017 that number spiked to 2,070 suspicious order reports.⁵⁰⁸ From 2005 to 2016, Cardinal Health distributed approximately 366 million dosage units of hydrocodone and oxycodone to its West Virginia customers, of which approximately 174 million dosage units or 47.5% were delivered between 2006 and 2011, the period where zero suspicious order reports could be confirmed. However, just doing the math, 52.5% or 192 million dosage units were delivered between 2011 and 2017 indicating that even with Cardinal reporting suspicious orders to the DEA, the company's efforts had little or no effect on actual amounts of opioids being supplied to West Virginia.

Shipment patterns in Summit and Cuyahoga counties also are illustrative. Beginning in approximately [REDACTED], Cardinal consistently shipped at least [REDACTED] dosage units per month of oxycodone and hydrocodone into Cuyahoga County.⁵⁰⁹ From [REDACTED] to a maximum of approximately [REDACTED] dosage units monthly.⁵¹⁰ Although that has [REDACTED], the level has [REDACTED].⁵¹¹

The case of Summit County is more dramatic. From [REDACTED], Cardinal shipped [REDACTED] dosage units per month of oxycodone and hydrocodone.⁵¹² Like Cuyahoga County, since [REDACTED] units per month.⁵¹³

CareMed Pharmacy

The history surrounding CareMed Pharmacy was explored during Mr. Reardon's deposition. The documentary evidence shown to Mr. Reardon demonstrated that the pharmacy owner in 2008 informed Cardinal's Quality and Regulatory Affairs personnel (Messrs. Morse and Forst) through the new pharmacy questionnaire that CareMed purchased opioids from multiple distributors including Cardinal.⁵¹⁴ Despite Mr. Forst acknowledging

⁵⁰⁸ See W.Va. Red Flags Report at 244.

⁵⁰⁹ See Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 51, 136 (Total Dosage of Oxycodone and Hydrocodone Shipped by Cardinal Health to Cuyahoga County, Ohio (Jan. 1996 to April 2018)).

⁵¹⁰ *Id.*

⁵¹¹ *Id.*

⁵¹² See Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 93, 178 (Total Dosage of Oxycodone and Hydrocodone Shipped by Cardinal Health to Summit County, Ohio (Jan. 1996 to April 2018)).

⁵¹³ *Id.*

⁵¹⁴ See S. Reardon Deposition at 404:21-24.

that he was aware of CareMed's use of multiple distributors, he allowed Cardinal to continue supplying CareMed resulting in a 609% increase in controlled substances purchasing over 13 months.⁵¹⁵

CVS #219

According to the DEA, from January 2008 to December 31, 2011, Cardinal sold over [REDACTED] dosage units of oxycodone to CVS #219 or on average [REDACTED] dosage units per month.⁵¹⁶ However, it appears that no on-site visit was conducted by Cardinal prior to February 2012.⁵¹⁷

From June 2009 to September 30, 2010, CVS #219 routinely exceeded its monthly oxycodone thresholds in amounts ranging from [REDACTED] dosage units in June 2009 to [REDACTED] units in August 2010.⁵¹⁸ At the same time, Cardinal approved threshold increases from [REDACTED] dosage units in June 2009 to a maximum of [REDACTED] dosage units per month from September 2010 with the notation that Cardinal did not see these amounts as unreasonable as to quantity, pattern or frequency.⁵¹⁹ However, Cardinal had clear evidence to the contrary.

In January 2010, Jennifer Hug, Manager of Retail National Accounts, wrote in an email that she "[s]poke with Nick at the pharmacy [and] [h]e stated that there are an increasing number of patients from pain clinics, but they are filing only for local clinics."⁵²⁰ Upon receiving a response from Jason Spinard in CVS's Loss Prevention Department that CVS #219 was okay, Ms. Hug forwarded the information to Christopher Forst and Maranda Swedyk and Michael Moné saying "I wanted to forward to you the information from CVS's LP department, in reference to the **2 SOM Events** attached."⁵²¹ Ms. Hug recognized that the orders from CVS #219 were suspicious, but Cardinal continued shipping oxycodone to the store anyway.

On September 30, 2010, Mr. Moné received an email from Paul Farley about Mr. Farley's conversation with CVS about CVS #219, noting that the increased sales of oxycodone was the result of store closures in the area and that as a result of the DEA "cracking down on 'pill mills' ... that is driving more legitimate traffic to CVS stores."⁵²² Mr. Farley also reported that CVS would not provide the doctor or patient information requested by Mr. Moné and that CVS did "not expect Cardinal to interrupt service to CVS stores since they have responded in the manner we agreed upon when launching the SOM program."⁵²³ Despite the weakness of the response and refusal to provide information, Mr. Moné approved the CVS orders on hold because "[w]e will be working

⁵¹⁵ See S. Reardon Deposition at 405:5-16.

⁵¹⁶ See Lakeland 2012 ISO at 2.

⁵¹⁷ See S. Morse Deposition at 117:13-17. (Morse although in charge of investigations and site visits did not know if a visit was conducted on CVS #219).

⁵¹⁸ See Cardinal Health Lakeland Thresholds Exceeded, CVS 219, CAH_MDL_PRIORPROD_DEA12_00004353.

⁵¹⁹ *Id.*

⁵²⁰ See Email from J. Hug to J. Spinard, CVS 850 & CVS 219 – LP Research (Jan. 27, 2010), CAH_MDL_PRIORPROD_DEA12_00011836.

⁵²¹ See Email from J. Hug to M. Swedyk, C. Forst, M. Moné, FW: CVS 850 & CVS – LP Research (Feb. 15, 2010) (emphasis added), CAH_MDL_PRIORPROD_DEA12_00011836.

⁵²² See Email from P. Farley to M. Moné, CVS #0219 (Sept. 30, 2010) (B. Jackson, SVP of National Accounts/Alternative Care was also copied on the email); CAH_MDL_PRIORPROD_DEA12_00003244 at CAH_MDL_PRIORPROD_DEA12_00003250.

⁵²³ *Id.*

through another solution.”⁵²⁴ At the same the held orders were released, Cardinal increased CVS #219’s oxycodone threshold by [REDACTED] (from [REDACTED] dosage units to [REDACTED] dosage units).⁵²⁵ Finally, on November 10, 2011, Mr. Moné touted that Cardinal reduced the oxycodone threshold for CVS #219 to [REDACTED] dosage units per month, but neglected to put that into the context that Cardinal had already sold [REDACTED] dosage units to the store.⁵²⁶

The results of this pattern of systemic failure in Cardinal’s anti-diversion program were entirely predictable. The amounts of opioids purchased, both legitimate and illegitimate, increased unabated for customers. Few, if any opioid orders, were deemed to be suspicious, investigated and reported to the DEA. Potential indicators that diversion was occurring were ignored.

10.4 Company Commitment – Compliance Culture, Organization & Resources

The evidence reviewed in this section demonstrates that Cardinal Health was and appears still not to be fully committed to meeting its regulatory obligations under the CSA and its accompanying regulations. As demonstrated by the culture, organizational structure and resourcing, compliance at Cardinal Health is secondary to management’s focus on driving increasing revenues and cutting costs.

10.4.1 Cardinal’s culture is myopically focused on increasing revenues and cutting cost while downplaying the importance of ethics and compliance.

In general, the Cardinal culture was and continues to be myopically focused on increasing revenues and cutting costs. Cardinal’s new CEO, Michael Kaufman, wrote in his FY 18 annual report letter to shareholders that a 2019 fiscal priority was to be “laser-focused on our cost structure” and “[s]imply put, we are intensely focused on each dollar we spend.”⁵²⁷ He also stated that the company is focused on “[a] corporate culture that drives success,” but what that translates into are “employees who are dedicated, hard-working and focused on the needs of their patients,” in other words a “high-performing, accountable culture.”⁵²⁸ “We roll up our sleeves to get the job done [and] [w]e know how to pull together in good times and bad.”⁵²⁹ Compliance, ethics, and integrity are not found in Mr. Kaufman’s stated vision of Cardinal culture. Instead, Cardinal Health’s culture is all about the bottom line.

This lack of putting compliance first carries over into Cardinal’s stated values. Cardinal values are:

1. We are **tenacious** in fulfilling our commitments to customers.

⁵²⁴ See Email from M. Moné to C. Forst (Sept. 30, 2010), CAH_MDL_PRIORPROD_DEA12_00003244 at CAH_MDL_PRIORPROD_DEA12_00003250.

⁵²⁵ See Cardinal Health Lakeland Thresholds Exceeded, CVS 219, CAH_MDL_PRIORPROD_DEA12_00004353.

⁵²⁶ See Moné Declaration at ¶ 45.

⁵²⁷ See Cardinal Health, *2018 Annual Report*, 3 (Sept. 2018).

⁵²⁸ *Id.* at 4.

⁵²⁹ *Id.*

2. We are **accountable** for high performance and to each other.
3. We are **inventive** and **adaptable**.
4. We bring a sense of **optimism, enthusiasm** and a **competitive spirit** to our work.
5. We are **inclusive** and **work together** with confidence and trust.
6. We are **genuine, open, direct** and **respectful**.
7. We can be **trusted** to do the right thing.⁵³⁰

Compliance is not among the listed Cardinal values, and when it is mentioned in the Standards of Business Conduct, Cardinal frames acting “with integrity and in compliance with the law,” within the frame of “[w]e work together, according to shared standards and values, to make wise decisions that foster a culture of trust and responsible conduct.”⁵³¹ This suggests that Cardinal only complies with a law where Cardinal shares the same standards and values as the law, but not does not necessarily comply when the company disagrees with it. In short, compliance with legal requirements is optional.

This cultural lack of commitment to compliance appears to be historically consistent within Cardinal. For example, in 2008, then Cardinal CEO, Kerry Clark sent an email to all Cardinal Global Leadership Team (“GLT”) members entitled “In the Penalty Box” to share some thoughts on regulatory compliance because “some of our failures in this area are burdening our financial performance and market capitalization.”⁵³² Mr. Clark went on to formulate his theory that “we keep getting penalties-big penalties that are costing us customer loyalty, employee loyalty, and shareholder loyalty. Big penalties that are undermining our progress on the path to premier [sic.] against all three stakeholder groups.”⁵³³ He saw the weakness in the Cardinal culture as a lack of “accountability of leadership” that resulted in employees who “do not fully trust their leaders.” Finally, he noted that “general managers are ultimately responsible for results [and] ... are accountable for ensuring their units operate according to quality and legal/regulatory standards.”⁵³⁴

However, that message of managerial accountability apparently did not take hold. In 2013, the McKesson team noted that “[p]erhaps the most surprising revelation [at the HDMA conference the prior week] was Steve Reardon and Gilberto Quintero saying Cardinal does not report suspicious orders to DEA ... no upside.”⁵³⁵ Mr. Reardon was Cardinal’s Vice President of Regulatory Operations, while Mr. Quintero was the Senior Vice President of Quality and Regulatory Affairs at Cardinal Health. Both were involved with overseeing and operating Cardinal’s controlled substances compliance program.

⁵³⁰ See Cardinal Health, *Standards of Business Conduct*, 2 (2018) (emphasis in the original) at <https://www.cardinalhealth.com/content/dam/corp/web/documents/fact-sheet/cardinal-health-standards-of-business-conduct-booklet-english.pdf>.

⁵³¹ *Id.* at 6.

⁵³² See Email from K. Clark to GLT Communication, In the Penalty Box (Jan. 17, 2008), CAH_MDL_PRIORPROD_DEA07_00827893.

⁵³³ *Id.*

⁵³⁴ *Id.*

⁵³⁵ See Email from W. de Guterrez-Mahoney I to D. Walker, *et al.*, HDMA notes (Mar. 11, 2013), MCKMDL00545341.

Based on the information I reviewed, it is readily apparent that Cardinal historically does not focus on or consider compliance with its controlled substances regulatory obligations particularly important, especially if this gets in the way of focusing on every dollar to maximize profits.

10.4.2 Cardinal failed to design an organizational structure with strong governance, made poor staffing choices and was slow to add resources to the SOM program demonstrating a lack of commitment to compliance.

During the review period, Cardinal has changed the structure of its controlled substances program several times. These changes correspond to Cardinal's various settlements with the government. However, while the Cardinal controlled substances program at times appeared robust on paper, in reality, it suffered from at least three underlying flaws: (a) a lack of strong senior management governance, (b) a slow commitment of resources and (c) poor choices of key operational staff.

Prior to 2007, Cardinal's central SOM organization was limited to three people, a vice president, and two managers.⁵³⁶ The Vice President of Quality and Regulatory Affairs, Steve Reardon, reported directly to the President and CEO of PDPS, Mark Parrish.⁵³⁷ In FY 2007 Compliance requested a 58% budget increase over FY2006 to add two FTE's (a manager and director).⁵³⁸ In presenting the business case for the two new positions, Compliance told senior management that "[c]urrent department staff workloads are at full capacity, [e]ffective management of current projects and initiatives is difficult[, and] [r]esources to take on new initiatives and the ability to improve and enhance existing programs are lacking."⁵³⁹ As Stephen Reardon Cardinal's Vice President Quality & Regulatory Affairs admitted in testimony, his staff could not investigate the volume of suspicious orders being identified.⁵⁴⁰

Starting in December 2007, with the approval of the two new positions, Cardinal slowly began increasing compliance resources. However, as described by Steve Lawrence the situation remained dire:

We currently are working very hard to staff up our QRA group. Please understand that they are working day, night and weekends but they have been understaffed.⁵⁴¹

Cardinal hired Craig Morford in 2008 to be its Chief Compliance Officer "with a mandate to establish a premier anti-diversion system."⁵⁴² On paper having the controlled substances program report to the Chief Compliance

⁵³⁶ See Board of Directors of Cardinal Health, Inc., *Investigation Report of the Special Demand Committee*, 7 (Apr. 12, 2013), CAH_MDL_PRIORPROD_HOUSE_0003331 ["Special Committee Report"].

⁵³⁷ See Drug Distribution, *Compliance Budget Review Fiscal Year 2007*, 5 (updated), CAH_MDL2804_02102331 at CAH_MDL2804_02102334 ["FY 2007 Budget"].

⁵³⁸ *Id.* at 3.

⁵³⁹ *Id.* at 2.

⁵⁴⁰ See Stephen Reardon Deposition, 466:23-470:21 (Nov. 30, 2018).

⁵⁴¹ See Steve Lawrence email to G-NSA-Regional Directors, *et al.*, FW: *Threshold system and customer issues – Detailed summary so I apologize for the lengthy email*, 1 (Jan. 26, 2008), CAH_MDL_PRIORPROD_DEA07_00891487.

⁵⁴² See Special Committee Report at 9.

Officer, who in turn reports to the CEO gives the appearance that Cardinal took controlled substances compliance seriously. However, that appearance diverges from the reality of the inner workings at Cardinal. Beginning in 2009, Mr. Morford's attention was split as he became Chief Legal Officer as well.⁵⁴³ Thus Morford was only fully engaged in establishing "a premier anti-diversion system" for 12 months. Mr. Morford's oversight role over the controlled substances program was further diminished with the hiring of Gilberto Quintero as Senior Vice President, Quality and Regulatory Affairs, who reported to Mr. Morford. While Mr. Quintero's hiring was necessary given the breadth of Mr. Morford's responsibilities, it is a further indication that Cardinal senior management provided tangential rather than direct oversight to the controlled substances program.

From 2008 to 2012, operational responsibility for the controlled substances program was vested in Michael Moné, Vice President, Anti-Diversion.⁵⁴⁴ Mr. Moné in early 2008 hired Steve Morse and Christopher Forst as Directors of Supply Chain Integrity.⁵⁴⁵ Their duties were split as follows:

- Morse was responsible for conducting investigations and site visits of customers, and
- Forst would handle the pharmaceutical analysis of customers.⁵⁴⁶

Therefore, in 2008, the central operational team for controlled substances compliance was still only 3 FTEs.

Cardinal very slowly increased headcount from 2010 to 2012. In 2010, Nicholas Rausch was added to the group as Director of Analytics and Information Management⁵⁴⁷ By 2012, the central anti-diversion team consisted of 8 FTEs.⁵⁴⁸

As of 2012, Messrs. Moné and Morse were no longer involved with the anti-diversion team. According to the Special Committee, Mr. Moné was replaced because "evaluation[s] of customers and orders had been heavily focused on the clinical expertise and subjective judgment of the pharmacists in the anti-diversion group," while Mr. Morse was removed because he was "not as strategic as his former position required and there were questions about his judgment."⁵⁴⁹ The removals of Messrs. Moné and Morse also coincide with Cardinal's 2012 AMOA.

From both an organizational design, as well as a resourcing perspective, Cardinal Health's failure to design an organizational structure with strong corporate governance together with its poor choice of staff and its slow commitment to add resources are consistent with the company's lack of focus on the importance of controlled substances compliance.

⁵⁴³ *Id.* at 9. According to Cardinal's official biography of Morford, his duties include responsibility for "Legal, Quality and Regulatory Affairs, Ethics and Compliance, Corporate Communications and Government Relations" *See* <https://www.cardinalhealth.com/en/about-us/our-people/our-leaders/craig-morford.html>.

⁵⁴⁴ *Id.* at 8

⁵⁴⁵ *Id.* at 8.

⁵⁴⁶ *See id.* at 8-9.

⁵⁴⁷ *Id.* at 9.

⁵⁴⁸ *See* Organization Chart 2012-2015, P1.4592.

⁵⁴⁹ *See* Special Committee Report at 34-35.

10.5 Program Core – Requirements, Education, Detection & Corrections

10.5.1 Viewed holistically, the core of Cardinal's SOM Program during the period ranges from incomplete to convoluted and inconsistent.

Overall the core of Cardinal's SOM program over the review period was either incomplete as it was before 2007, or too convoluted and inconsistent thereafter. In either case, it demonstrates that Cardinal failed to implement a program that was consistent with its understanding of its controlled substances obligations.

For the program pre-2007, the primary issue is that Cardinal artificially limited the focus of the program by concentrating only on reporting information to the DEA, rather than on its statutory obligation to prevent the product from being diverted. As a result, the pre-2007 system, as designed and implemented, was incomplete.

Cardinal's program from 2007 to 2012 was convoluted in that it was difficult to determine across the five key SOPs that comprised the program, where one ends, and the other begins.⁵⁵⁰

For example, it is unclear from the SOPs how the On-site Investigations and Detecting and Reporting Suspicious Orders and Responding to Threshold Events procedures operate together to form the suspicious order due diligence component of the program.

Additionally, some key definitions that comprise Cardinal's SOM program are not consistent across the various SOP's even though they are part of the same series (CAD-Cxxx). For example, the definition of "threshold event" (see table below) varies widely across the SOPs.

Table 1 Varying Definitions of "Threshold Event"

SOP	YEAR	DEFINITION
ON-SITE INVESTIGATIONS⁵⁵¹	2009	An order for a regulated drug which exceeds the threshold set for a specific licensed customer.
SALES - ANTI-DIVERSION ALERT SIGNALS⁵⁵²	2009	Threshold Event Is defined as the initial held order created by a DEA#, Base Code, Threshold Limit combination.

⁵⁵⁰ The four key SOP's are: Process to Establish SOM Threshold Limits, Sales-Anti-Diversion Alert Signals, On-site Investigations, and Detecting and Reporting Suspicious Orders and Responding to Threshold Events. See Cardinal Health, Process to Establish SOM Threshold Limits, HSCSQRA-CAD-C-002 (Dec. 22, 2008), CAH_MDL_PRIORPROD_AG_0004208. In 2010 as the result of a restructuring, this document was renumbered as PDQRA-CAD-C002 via Document Control Notice ("DCN") 2555 (CAH_MDL_PRIORPROD_AG_0000017); Cardinal Health, Sales-Anti-Diversion Alert Signals, HSCSQRA-SAD-C003 (Dec. 22, 2008), CAH_MDL_PRIORPROD_AG_0000323; Cardinal Health, On-site Investigations, PDQRA-CAD-C008 (Nov. 5, 2009), CAH_MDL_PRIORPROD_DEA12_00014535 ["Investigations SOP"]; Cardinal Health, Detecting and Reporting Suspicious Orders and Responding to Threshold Events, PDQRA-CAD-C007 (Jan. 6, 2012), CAH_MDL_PRIORPROD_AG_0000154 ["Reporting SOs 2012 SOP"]; Cardinal Health, Large Volume- Tactical and Analytical Committee Periodic Review Process, PRDQRA-CAD-C023 (Apr. 12, 2012), CAH_MDL2804_02288612 ["LV-TAC SOP"].

⁵⁵¹ See Investigations SOP at § 4.1.

⁵⁵² See Sales Alert SOP 2009 at § 4.1.

SOP	YEAR	DEFINITION
DETECTING AND REPORTING SUSPICIOUS ORDERS AND RESPONDING TO THRESHOLD EVENTS ⁵⁵³	2012	The initial held order for a regulated drug which exceeds the threshold set for the specified customer. This is created by a DEA#, Base Code and Threshold Limit combination.

10.5.2 Cardinal Health failed to understand that it and not the DEA was responsible for maintaining an effective program to detect and report suspicious orders and prevent diversion.

In 2012, Michael Moné, Vice President of Supply Chain Integrity, made a declaration that Cardinal repeatedly requested the DEA supply Cardinal with any information it had on Cardinal customers engaging in diversion.⁵⁵⁴ For example, in October 2011, Cardinal's Chief Legal Officer, Craig Morford, asked DEA to provide that information and "Cardinal Health promised to immediately cease distributing controlled substances to any customer DEA so identified."⁵⁵⁵ Mr. Morford stressed the issues of DEA's "[l]ess engagement with the industry," and "[l]imited guidance or notice" to Cardinal's Board of Directors Audit Committee meeting in November 2012.⁵⁵⁶

The declarations by Messrs. Morford and Moné demonstrate that Cardinal fundamentally misunderstood its obligations under the CSA. First, the CSA places an affirmative duty on distributors of controlled substances to maintain an effective program to prevent diversion.⁵⁵⁷ Second, the CSA places no affirmative duty on the DEA to disclose any information to Cardinal regarding Cardinal customers engaging in diversion. Third, Mr. Moné's statement and Mr. Morford's presentation imply that because the DEA provided no information to Cardinal to "indicate that any of its customers was [sic.] diverting controlled substances," that diversion was not occurring.⁵⁵⁸ Based on the evidence readily available to Cardinal, this was not the case.

10.5.3 Cardinal's early controlled substances program was not compliant with DEA regulatory requirements.

Cardinal's Board of Directors in November 2012 formed a "Special Demand Committee" to respond to a shareholder demand letter alleging that as evidenced by the 2012 AMOA, Cardinal failed to implement and

⁵⁵³ See Reporting SOs 2012 SOP at § 4.1.

⁵⁵⁴ See Cardinal Health, Inc. v. Eric Holder, Jr., et al, Declaration of Michael A. Moné Pursuant to 28 U.S.C. § 1746 (Feb. 3, 2012) at ¶ 35, CAH_MDL_PRIORPROD_DEA12_00014053 at CAH_MDL_PRIORPROD_DEA12_00014067-68 [Moné Declaration]

⁵⁵⁵ See *id.*

⁵⁵⁶ See Presentation by C. Morford, 2012 Annual Quality and Regulatory (QRA) Report to the Audit Committee of the Board of Directors, 8 (Nov. 2, 2012), CAH_MDL2804_03262274 at CAH_MDL2804_03262440.

⁵⁵⁷ See generally Section 5, *infra*.

⁵⁵⁸ See Moné Declaration at ¶ 35.

maintain systems to detect and prevent diversion as required under the CSA.⁵⁵⁹ The Special Committee reported that prior to 2007, Cardinal focused its anti-diversion or AD measures on preventing price diversion and internet pharmacy diversion.⁵⁶⁰ The Special Committee also noted that the controlled substances program had only a few employees dedicated to it, had no electronic order analyzer system, therefore, most of the customer and order information was paper-based and scattered.⁵⁶¹ Cardinal told a similar story to the House Energy & Commerce Committee (“E&C Committee”) in 2018 and specifically acknowledged it was not until December 2008 that “Cardinal implemented formal anti-diversion Standard Operating Procedures (SOPs), which included SOPs for conducting prospective customer due diligence.”⁵⁶²

Cardinal issued a Corporate Quality & Regulatory Compliance Standard Operating Procedures manual in June 2006 under the approval of Stephen Reardon, Cardinal’s Vice President Quality & Regulatory Affairs.⁵⁶³ At that time, the primary internal control was the use of the Ingredient Limit Report. The report, which was provided to the DEA on a monthly basis, identified customers that exceeded a pre-determined purchase level.⁵⁶⁴ According to Mr. Reardon, the Ingredient Limit Report was used by Cardinal from approximately 1994 to 2008.⁵⁶⁵

The SOP manual described the Ingredient Limit Reports as “based on a computer program which monitors customer controlled substance purchases for a month and compares these purchases to predetermined averages or limits and if a customer’s purchase quantities exceed the established parameters, the customer’s activity is printed on the report.”⁵⁶⁶ However, the Ingredient Limit Reports only identified orders exceeding limits. They did not prevent the identified orders from shipping.⁵⁶⁷

In testimony to the E&C Committee, Cardinal stated that “[t]he reports were generated based on a computer algorithm established by the DEA, which was meant to be used to calculate the quantity which, if exceeded in one month, constituted an order which may be excessive or suspicious.”⁵⁶⁸ Despite these descriptions, the methodology on how the limits were developed in Ingredient Limit Reports remains exceedingly unclear.

⁵⁵⁹ See Special Committee Report at 4-5.

⁵⁶⁰ *Id.* at 7.

⁵⁶¹ *Id.* at 8.

⁵⁶² See W.Va. Red Flags Report at 115.

⁵⁶³ See Cardinal Health, *Corporate Quality & Regulatory Compliance Standard Operating Procedures*, 1 (Jun. 15, 2006), CAH_MDL_PRIORPROD_DEA07_01188323 [“CAH 2006 SOP Manual”]. Prior to this manual, Cardinal had issued the DEA Compliance Manual a collection of documents that was “intended as a resource to the Controlled Substances Act and Regulations ... [and] as a guide so that you can better understand DEA’s function. Further, it is intended to help you learn to deal with the agency that has a tremendous impact on our business.” See Cardinal Health, *DEA Compliance Manual* CAH_MDL_PRIORPROD_DEA07_01383895 at CAH_MDL_PRIORPROD_DEA07_01383902.

⁵⁶⁴ See Special Committee Report at 8.

⁵⁶⁵ See S. Reardon Deposition at 411:20-23.

⁵⁶⁶ See CAH 2006 SOP Manual, *DEA04.00 - Required Reports to DEA*, at 6 § 5(c).

⁵⁶⁷ See S. Reardon Deposition at 427:12-23.

⁵⁶⁸ See W.Va. Red Flags Report at 184.

However, according to Mr. Reardon, the limits were a combination of averages and multipliers to reach the final threshold limits.⁵⁶⁹

According to the SOP Manual, the distribution facility was required to monitor and identify “individual orders that appear excessive in relation to what other customers are buying and/or the customer's purchase history” and “to notify the local DEA field office, if possible before the order [was] shipped.”⁵⁷⁰ Mr. Reardon testified that from 1994 to 2008 Cardinal relied on the “pickers and checkers” filling orders in the distribution center to identify the individual orders.⁵⁷¹

Thus, DEA notification was the main purpose of Cardinal's SOM system prior to 2007. The program at that time did not require Cardinal employees to actively prevent suspicious orders from shipping. Employees merely needed to try to notify the DEA before the order shipped.

Therefore, as set out in the SOP Manual, Cardinal's controlled substances compliance program was not compliant with the DEA requirements as Cardinal employees were not expressly required to investigate “suspicious” orders or stop suspicious shipments to avoid potential diversion. Simply using thresholds to provide notices to the DEA does not constitute an operational program to prevent diversion. Thus, Cardinal's pre-2007 controlled substances compliance program efforts were not compliant as designed.

Cardinal's reliance on Ingredient Limit Reports and distribution center pickers did not work.⁵⁷² The chart below from DEA's Show Cause Order for Lakeland, Florida bears this out. The magnitude of the failure is especially egregious when contrasted against the fact that according to the DEA most retail pharmacists in Florida at that time were ordering less than 8,400 dosage units per month of hydrocodone.⁵⁷³

⁵⁶⁹ See S. Reardon Deposition at 442:12-16.

⁵⁷⁰ See CAH 2006 SOP Manual at 6 § 5(d)(i).

⁵⁷¹ See S. Reardon Deposition at 428:10-24 and 429:1-2.

⁵⁷² See Lakeland 2007 ISO at 2.

⁵⁷³ *Id.*

Table 2: Chart of Cardinal Florida Customers DEA Believed Were Diverting Hydrocodone⁵⁷⁴

Pharmacy	Total Dosage Units	Number of Months Distributions Made	Monthly Average	Dates of Distribution (*Distributions not made in every month)
Medipharma-Rx, Inc.	620,030	4	155,007	Aug – Dec 05*
DRM Enterprises, Inc.	929,600	22	42,254	Jan 06 – Oct 07
Jen-Mar Pharmacy Services, Inc.	353,700	11	1 st 3 mos: 32,154 Last 8 mos: 2,766 43,175	Mar 06 – Feb 07*
Armenia Pharmacy, Inc.	132,900	12	1 st 6 mos: 11,075 Last 6 mos: 1,900 20,250	Mar 06 – Feb 07
National Pharmacy, Inc.	659,800	9	73,311	Aug 05 – May 06*
Parulmed Corporation	468,400	20	23,420	Aug 05 – Apr 07*
Q-R-G, Inc.	1,213,200	5	242,640	Feb – June 06
RKR Holdings, Inc.	741,000	13	57,000	Aug 05 – Jan 07*
United Prescription Services, Inc.	1,148,100	4	287,025	Jul – Oct 06
Satellite Drug and Pharmacy	1,044,000	19	1 st 4 mos: 54,947 Last 15 mos: 375 69,500	Feb 06 – Oct 07*

10.5.4 Cardinal's early training efforts were not fit for compliance purposes.

The Special Committee report described Cardinal's efforts to educate employees about controlled substances in the following terms:

During Spring and Summer 2006, there were live training sessions for employees at six locations around the country, focused primarily on price diversion and diversion by internet pharmacies. The Company also developed and implemented two mandatory computer-based trainings in August and December 2007 for all sales personnel and field operations managers, which also focused on internet pharmacy diversion.⁵⁷⁵

As described, Cardinal's training efforts were defective in at least three regards. First, the training, like the rest of the controlled substances program at that time, was inappropriately limited to price diversion and internet pharmacies rather than being a holistic anti-diversion program for controlled substances. For example, in 2007 Know Your Customer training, Eric Brantley, Director of Quality and Regulatory Affairs, told participants that

⁵⁷⁴ See Lakeland 2007 ISO at 2.

⁵⁷⁵ See Special Committee Report at 11.

questionnaires were being used to look for red flags, which he defined as “[l]ooking for signs of Internet activity (shipping supplies, lack of walk-in customers, etc.) and filling prescriptions from questionable ‘pain clinics’.”⁵⁷⁶ This is an artificially narrow, and inaccurate, rendition of the DEA’s requirements pertaining to suspicious orders.

Second, before the training efforts described above, the Special Committee report did not reference any earlier substantial controlled substances training. This suggests that prior to 2006 Cardinal had no concerted training effort even though the CSA and the implementing regulations were in effect since 1971. A January 2005 Quality management meeting presentation provided confirmation on this point as the Enterprise Training Initiative section noted:

- There was a lack of corporate sponsorship for training;
- Training was left to the individual sites and was non-existent at some sites;
- There were no repercussions for not completing training or failing assessments; and
- There was both insufficient and redundant training.⁵⁷⁷

This lack of earlier training suggests that Cardinal’s efforts in 2006-2007 appear to be more of an attempt by the company to demonstrate the existence of a controlled substances compliance program in the face of mounting enforcement pressures, rather than a genuine corporate effort to discharge its obligation to maintain an effective program to prevent diversion.

Third, the Special Committee specifically referenced that Cardinal developed two mandatory computer-based training courses but did not describe the content or note whether participants were required to pass an assessment to demonstrate comprehension of the information. While highlighting the number of courses given and the number of employees trained, as Cardinal does, are common compliance training metrics, they only are measures of activity rather than effectiveness. Therefore, the metrics cited by the Special Committee are useless in determining whether compliance training activities were effective.

10.5.5 Cardinal’s SOM program modified in response to the DEA enforcement proceedings failed to meet the necessary standards in five key areas: threshold setting, threshold increases, due diligence, customer monitoring, and investigations.

As a result of the 2007 ISOs, Cardinal undertook a series of program modifications in order to reach the 2008 settlement agreement. These changes included adding additional headcount and adjusting the organizational structure, implementing new customer due diligence requirements, issuing new standard operating procedures, implementing an electronic monitoring system, and modifying how Cardinal set and utilized thresholds.⁵⁷⁸ The Cardinal controlled substances program in this timeframe is defined largely by five SOPs:

⁵⁷⁶ See E. Brantley presentation, *Know Your Customer Retail Pharmacy Questionnaire Training*, 12 (Oct. 2007), CAH_MDL_PRIORPROD_DEA07_02214751. [“KYC 2007 Training”]

⁵⁷⁷ Operation One Cardinal Health Quality Management Meeting, Enterprise Training Initiative Quality/Regulatory, 30 (Jan. 13, 2005), CAH_MDL_PRIORPROD_DEA07_01181262 at CAH_MDL_PRIORPROD_DEA07_01181276.

⁵⁷⁸ See Special Committee Report at 9-11.

- Process to Establish SOM Threshold Limits,⁵⁷⁹
- Sales-Anti-Diversion Alert Signals,⁵⁸⁰
- On-site Investigations,⁵⁸¹
- Detecting and Reporting Suspicious Orders and Responding to Threshold Events and⁵⁸²
- Large Volume- Tactical and Analytical Committee Periodic Review Process.⁵⁸³

However, as discussed below, the program modifications undertaken did not result in an effective controlled substances compliance program. This occurred in part because the design and the actual SOPs were ambiguous, creating opportunity to circumvent the controls and because Cardinal personnel did not follow the established processes.

A. Establishing Thresholds

The December 2008 SOP entitled “Process to Establish SOM Threshold Limits” outlined how thresholds were set. The methodology defined in the SOP was complex and contained several sub-steps, but in general, to establish a threshold Cardinal would:

(1) extract and formulate a list of customers that have purchased monitored items and historical sales data for those customers for all monitored items; (2) differentiate customers through segmentation by size and/or specialty; (3) evaluate historical controlled substance sales data per drug family, per month for each customer segment to establish appropriate threshold limits, using the multiples of 3, 5, or 8; (4) incorporate background information about the pharmacies to establish final threshold limits; and (5) apply rounding logic and finalize threshold limits.⁵⁸⁴

Cardinal told the House Committee that the new custom threshold limits were:

designed to alert analysts automatically whenever a customer’s order volume exceeded its assigned threshold. All orders that triggered threshold events were held and reviewed to determine whether the order was justified or was suspicious [and] [o]rders that were determined to be suspicious were not shipped.⁵⁸⁵

⁵⁷⁹ See Cardinal Health, Process to Establish SOM Threshold Limits, HSCSQRA-CAD-C-002 (Dec. 22, 2008), CAH_MDL_PRIORPROD_AG_0004208. In 2010 as the result of a restructuring, this document was renumbered as PDQRA-CAD-C002 via Document Control Notice (“DCN”) 2555 (CAH_MDL_PRIORPROD_AG_0000017).

⁵⁸⁰ See Cardinal Health, Sales-Anti-Diversion Alert Signals, HSCSQRA-SAD-C003 (Dec. 22, 2008), CAH_MDL_PRIORPROD_AG_0000323.

⁵⁸¹ See Cardinal Health, On-site Investigations, PDQRA-CAD-C008 (Nov. 5, 2009), CAH_MDL_PRIORPROD_DEA12_00014535 [“Investigations SOP”].

⁵⁸² See Cardinal Health, Detecting and Reporting Suspicious Orders and Responding to Threshold Events, PDQRA-CAD-C007 (Jan. 6, 2012), CAH_MDL_PRIORPROD_AG_0000154 [“Reporting SOs 2012 SOP”].

⁵⁸³ See LV-TAC SOP 2012.

⁵⁸⁴ See Special Committee Report at 12.

⁵⁸⁵ See W.Va. Red Flags Report at 185.

The threshold formula, however, like the Ingredient Limit Report levels that preceded it, remains a mystery. For example, the threshold formula requires that once a monthly limit is determined from historical sales data, if the product is hydrocodone or oxycodone, the monthly limit is to be multiplied by a factor of 3.⁵⁸⁶ The SOP presents no explanation or rationale for why hydrocodone or oxycodone monthly limits should be increased three-fold.⁵⁸⁷ The Special Committee report suggests that the multiples were derived from DEA guidance “to chemical distributors approving the use of multiples of three and eight for drug products consisting of a listed chemical and a controlled substance.”⁵⁸⁸ However, the citation provided in the Committee report could not be verified as the webpage no longer exists.

However, thresholds could be increased by more than the factor of three. According to the SOP, Cardinal could apply a final set of additional upwards adjustments based upon background or Know Your Customer (“KYC”) information.⁵⁸⁹ The stated reason for this provision was to “adjust threshold limits based on the associated level of risk,” such as increasing the threshold if the customer had “a documented diversion or loss prevention program.”⁵⁹⁰ This additional final adjustment, however, also provided Cardinal an opportunity to negate the algorithm and potentially make unsubstantiated and unregulated KYC adjustments resulting in threshold levels in excess of the system generated levels. Furthermore, the example provided in the SOP only requires a documented diversion or loss prevention program. Cardinal, therefore, was not required to ascertain whether the program worked in practice or was even being used. The net result of the KYC provision is that it creates a potentially significant “loophole” allowing Cardinal to adjust thresholds to satisfy customer needs regardless of the diversion potential.

In developing the new threshold methodology, Cardinal Health enlisted the help of several consultants, including:⁵⁹¹

- **Deloitte.** Cardinal Health retained Deloitte at various times between 2007 and 2012. Part of Deloitte's work was to assist in a project management capacity with business and technology enhancements to Cardinal Health's Controlled Substance Anti-Diversion program.
- **Dendrite/Cegedim/Buzzeo PDMA.** BuzzeoPDMA and Dendrite were both acquired by Cegedim, which is now part of IQVIA (formerly IMSQuintiles). Cardinal Health engaged Dendrite in 2007 to consult on Cardinal Health's on-site investigations program and used Dendrite to assist with on-site pharmacy investigations. Cardinal Health continues to use investigators from Buzzeo/Cegedim to assist with on-site pharmacy investigations.

⁵⁸⁶ See HSCSQRA-CAD-C002 at § 4.2.3(b)(vi).

⁵⁸⁷ It appears that the factor of 3 was derived from the Chemical Handler's Manual. Appendix E-3 entitled “Suspicious Order Reporting System for Use in Automated Tracking Systems.” See Discussion *infra* at Section 5.3.2.

⁵⁸⁸ See Special Committee Report at 12, n. 9.

⁵⁸⁹ See HSCSQRA-CAD-C002 at § 4.2.4

⁵⁹⁰ *Id.*

⁵⁹¹ See Cardinal Health, Inc., First Supplemental Objections and Responses to Plaintiffs' First Combined Discovery Requests at 13-14, *In Re: National Prescription Opiate Litigation*, Master Docket No.1:17-MD-02804-DAP MDL No. 2804 (N.D. Ohio).

However, Cardinal Health appears not to have been fully transparent when describing how it used those consultants. For example, the SOP provides that “[i]n the event that an adequate sample does not exist to formulate a threshold limit for a Base Code, initial threshold limits established for the segment by **Deloitte** will be used as a baseline.”⁵⁹² A January 2008 email from Carolyn McPherson, Director, Quality and Regulatory Affairs, was more explicit stating “Attached is [sic.] the thresholds by customer group list supplied by Deloitte.”⁵⁹³ Based on both the SOP and the email, Deloitte’s role was more significant than simple project management as Deloitte apparently developed at least some threshold limits on behalf of Cardinal.

In the same vein, it appears that Cardinal engaged Dendrite/Cegedim/BuzzeoPDMA to do more than just assist with on-site pharmacy investigations, and instead they did a full-blown review of Cardinal’s SOM system.⁵⁹⁴ That engagement included “[a]n onsite review of Cardinal’s suspicious order monitoring program/system including verification of the system’s operational effectiveness, system integrity, and **regulatory suitability**.”⁵⁹⁵ This verification of regulatory suitability is not consistent with on-site pharmacy investigations.

What is most concerning is that while this type of misrepresentation or dissembling may be couched as a “litigation strategy,” it speaks directly to Cardinal’s culture and a lack of integrity; once more reinforcing the conclusion that Cardinal does not take its ethical and compliance obligations seriously.

B. Threshold Events

During this period Cardinal also implemented the Detecting and Reporting Suspicious Orders and Responding to Threshold Events SOP. Although ultimately retired in 2016,⁵⁹⁶ the purpose of the SOP was to comply with or “exceed” distributor standards in CSA and DEA requirements.⁵⁹⁷ The procedure also was intended to:

provide guidance to Cardinal Health (CAH) employees in the Quality and Regulatory Affairs (QRA) section on responding detecting and reporting suspicious orders, and processing, documenting and making judgments about threshold events, including making decisions about releasing or cutting orders that are suspicious or exceed a threshold.⁵⁹⁸

The procedure required QRA to review every order that was a held or cut (removal of the offending line items).⁵⁹⁹ Held or cut orders were only deemed suspicious for DEA purposes if they meet at least one of three

⁵⁹² See HSCSQRA-CAD-C-002 at § 4.2.3(c).

⁵⁹³ See Email from Carolyn McPherson to S. Reardon, *et al.*, Deloitte Threshold Values By Type_Base_Size Combo-Report 2a.xls, 1 (Jan. 28, 2008), CAH_MDL_PRIORPROD_DEA07_00863981.

⁵⁹⁴ See S. Reardon Deposition at 460:11-463:15 (Discussing Cegedim’s work product and Reardon’s role in it).

⁵⁹⁵ See Email from Paul Hamby to J. Bennett, *et al.*, Cegedim Dendrite SOM review, 1 (December 15, 2007) (emphasis added) (Mr. Hamby was Cegedim’s Director of Consulting and Validation Services.), CAH_MDL_PRIORPROD_DEA07_00869802.

⁵⁹⁶ See Cardinal Health, Detecting and Reporting Suspicious Orders and Responding to Threshold Events, PDQRA-CAD-C007 (Oct. 17, 2016), CAH_MDL_PRIORPROD_AG_0001281 [“Reporting SOs 2016”].

⁵⁹⁷ See Reporting SOs 2012. at § 1.2.

⁵⁹⁸ See Reporting SOs 2012 at § 1.1.

⁵⁹⁹ *Id.* at § 6.1.3

criteria: (a) order is of unusual size, (b) order is of unusual frequency, and (c) order deviates substantially from a normal pattern for the customer.⁶⁰⁰

Orders of unusual size were defined as those “**significantly larger** than orders normally placed by the customer or by **customers that have a size and type of business that is similar** to the ordering customer’s business.”⁶⁰¹ Orders of unusual frequency were defined as those “orders that occur **significantly more frequently** than the orders normally placed by the ordering customer or by **customers that have a size and type of business that is similar** to the ordering customer’s business.”⁶⁰² Orders that deviate substantially from the normal ordering pattern were defined as “orders that reflect a **significant deviation** from the customer’s normal ordering pattern or that deviate substantially from the ordering patterns of **customers that have a size and type of business that is similar** to the ordering customers business.”⁶⁰³

Cardinal’s process however does not define “significantly larger,” “significantly more frequently,” or “significant deviation.” Therefore, it is unclear what significant means in this context. The same situation existed with “customers that have a size and of business that is similar.” Cardinal’s process failed to outline how a similar customer was to be determined for purposes of comparison. However, in all three cases, the SOP just required QRA personnel to use available information and experience to make reportability determinations.⁶⁰⁴

The SOP also notes that unusual size errors might be unintentional (e.g., typographical errors or duplicate orders). The SOP also was unusually vehement requiring that:

Unintentional order entry errors (including duplicate order entries) **MUST NOT** be reported as suspicious orders to DEA since the customer did not intend to place the order and **MUST** be cut with no changes to customer threshold and a readjustment of accrual to the level prior to the order entry error. (Emphasis in the original).⁶⁰⁵

Since reporting suspicious orders to the DEA did not guarantee regulatory action, it is highly usual for a company to worry about possible over-reporting. Furthermore, the emphasis suggests that Cardinal was looking for reasons not to report suspicious orders, which is reinforced by sudden removal of the language from the 2016 version of the SOP. Finally like the issues with the Alert Signals process, through the lack of precisely defined standards, Cardinal created a program that was susceptible to wide variation and being circumvented. Therefore, a claim that the program was an effective anti-diversion program is not supportable.

In another example of how Cardinal’s program is convoluted and difficult to follow, Cardinal in 2012 implemented the Large Volume – Tactical and Analytical Committee (“LV-TAC”) process which operated in

⁶⁰⁰ *Id.* at § 6.1.3(a-c).

⁶⁰¹ *Id.* at § 6.1.5 (emphasis added).

⁶⁰² *Id.* at § 6.1.6 (emphasis added).

⁶⁰³ *Id.* at § 6.1.7 (emphasis added).

⁶⁰⁴ *Id.* at §§ 6.1.5.1, 6.1.6.1 and 6.1.7.1. These sections remained unchanged through the 2016 version. *See* Reporting SOs 2016 at §§ 6.1.5, 6.1.6 and 6.1.7.

⁶⁰⁵ *Id.* at § 6.1.5.2.

parallel to the Detecting and Reporting Suspicious Orders and Responding to Threshold Events.⁶⁰⁶ The LV-TAC was a committee comprised of the Senior Vice President QRA, Regulatory, Vice President of Supply Chain Integrity and Director of QRA Analytics, and VP of Sales or appropriate designee all of whom met “on a periodic basis (e.g., monthly)” to assess customers via a detailed review of “the top retail purchasers of commonly diverted controlled substances.”⁶⁰⁷

It is not clear how Cardinal determined which customers were on the list. According to the SOP, “large purchasers of controlled substances” meant “[t]op retail purchasers of commonly diverted controlled substances,” but could be generated “for the entire network” or “by distribution center and will include all CAH customers classified as ‘Retail Pharmacies.’” Thus, when Cardinal defined the “top retail purchasers” it is unclear if that meant the top ten, top fifty or top one hundred. Furthermore, as with the other SOM procedures reviewed, Cardinal watered down the LV-TAC process in 2014 by eliminating the VP of Sales as a required committee member and removing the suggestion that periodic equaled monthly.⁶⁰⁸ However, Cardinal also stripped out any reference to how the Committee’s decisions go beyond the paper file and translate into action. The final step in the 2014 version of the process did not consist of notifying sales of the Committee’s decision and taking some action as in the 2012 version, but rather required Cardinal to simply collect and file the data reviewed by Committee into the digital filing system.⁶⁰⁹ Thus, by 2014, Cardinal seems to have eviscerated the Committee’s authority.

C. New Pharmacy Questionnaire

Cardinal Health told the House E&C Committee that:

In 2007, Cardinal Health began requiring completion of a New Pharmacy Questionnaire as part of the account approval process for all new retail independent pharmacies. The questionnaire collected general information about the pharmacy, its owner, and the pharmacist in charge; general information about the pharmacy’s other suppliers; information about the pharmacy’s customers and their primary method of payment for controlled and non-controlled substances; and the pharmacy’s expected controlled substance ordering, among other information. Cardinal Health employees vetted these questionnaires and conducted an additional investigation where appropriate.⁶¹⁰

As part of the 2008 settlement agreement, Cardinal agreed:

[t]o the extent it has not otherwise done so, Cardinal shall conduct an investigation for each customer where such review reveals purchasing patterns substantially deviating from the normal purchasing

⁶⁰⁶ See LV-TAC SOP 2012.

⁶⁰⁷ *Id.* at §§ 1.1 and 4.0.

⁶⁰⁸ See Cardinal Health, Large Volume- Tactical and Analytical Committee Periodic Review Process, PRDQRA-CAD-C023, §§ 1.1 and 4.0 (May 29, 2014), CAH_MDL_PRIORPROD_CNTY_0000450.

⁶⁰⁹ See LV-TAC SOP 2012 at § 7.0 *but compare* LV-TAC SOP 2014 at § 7.0.

⁶¹⁰ See W.Va. Red Flags Report at 115.

patterns, and take appropriate action as required by this Agreement, DEA regulations, and other procedures established under Cardinal's compliance program.⁶¹¹

In the October 2007 training program about the new questionnaire, Mr. Brantley told participants that the Pharmacy Business Consultant was responsible for completing and submitting the questionnaire to Corporate Quality and Regulatory Affairs for approval after working with pharmacy owner.⁶¹² He also made it clear that the Pharmacy Business Consultant was “the first line of defense for Cardinal in preventing diversion,” and therefore, “[t]he questionnaire must be taken seriously and must be filled out thoroughly and completely.”⁶¹³

Despite the admonitions in the October 2007 training and the agreement with the DEA in 2008, the DEA in 2012 alleged that Cardinal failed to conduct meaningful due diligence on “its retail pharmacy chain customers.”⁶¹⁴ Mr. Reardon in his deposition confirmed that Cardinal was relying on the chain customers to self-police and thus did not undertake independent due diligence using Cardinal employees.⁶¹⁵ Beginning in 2012, Cardinal changed its practices to stop relying simply on the chain store’s internal due diligence.⁶¹⁶

D. Anti-Diversion Alert Signals

At the same time Cardinal moved to individualized thresholds, the company also instituted new procedures to address customer monitoring and the handling of threshold excursions. In December 2008, Cardinal issued the “Sales – Anti-Diversion Alert Signals” SOP,⁶¹⁷ which was effective for only six months until it was completely rewritten “to conform to existing Cardinal Health practices.”⁶¹⁸

The purpose of the SOP was, as part of Cardinal’s overall SOM program, to provide “process requirements for the continuous monitoring and reporting of customer order activities by Sales during the execution of the SOM program.”⁶¹⁹ The SOP’s focus was on “threshold event investigation activities.”⁶²⁰

⁶¹¹ See 2008 CAH AMOA at 4.

⁶¹² See KYC 2007 Training at 10.

⁶¹³ *Id.*

⁶¹⁴ See Letter from M. Leonhart, Order to Show Cause and Immediate Suspension of Registration, Lakeland, Florida Distribution Center, 3 (Feb. 2, 2012), CAH_MDL2804_02465982 at CAH_MDL2804_02465995 [“Lakeland 2 ISO”].

⁶¹⁵ See S. Reardon Deposition at 401:21-402:21.

⁶¹⁶ See Presentation by C. Morford, *2012 Annual Quality and Regulatory (QRA) Report to the Audit Committee of the Board of Directors*, 8 (Nov. 2, 2012), CAH_MDL2804_03262274 at CAH_MDL2804_03262440 (“Large Chain Customers – can’t rely on controls of large, publicly traded chains; will conduct our own due diligence.”).

⁶¹⁷ See Cardinal Health, Sales Anti-Diversion Alert Signals (Dec. 22, 2008), CAH_MDL_PRIORPROD_AG_0000323 [“Sales Alert SOP 2008”].

⁶¹⁸ See Cardinal Health, Sales Anti-Diversion Alert Signals, HSCSQRA-SAD-C003 at Change History (Jun. 9, 2009) (Document was changed by DCN-2423), CAH_MDL_PRIORPROD_AG_0000326 [“Sales Alert SOP 2009”]; see also Cardinal Health, Sales Anti-Diversion Alert Signals, HSCSQRA-SAD-C003, (May 1, 2013) (Showing the SOP as being retired as of that date), CAH_MDL_PRIORPROD_HOUSE_0000264.

⁶¹⁹ See Sales Alert SOP 2008 at § 1.0.

⁶²⁰ *Id.*

According to the Alert Signals SOP, a “threshold event” was a situation where a customer’s order was held because the “customer’s accrual for a drug family in a given month surpasses the assigned threshold limit.”⁶²¹ According to the SOP, once the threshold event is triggered, the order is held “pending regulatory review.”⁶²²

The definition of “held order” was significantly changed between the 2008 and 2009 SOP versions regarding the treatment of subsequent orders and the handling of notifications. In the 2008 version, subsequent orders by the customer in the same drug family would be held “unless the threshold is increased by QRA, or unless the accrual is reset to zero at the beginning of the month.”⁶²³ The 2009 version simply provided that subsequent orders would be held “as a continuation of the original event.”⁶²⁴ Despite DCN-2423’s rationale for the change, this difference in wording appears to be one to remove potentially problematic wording rather than a change in practice. The 2009 version does not prevent QRA from increasing thresholds nor does it clearly address what happens when the monthly accrual resets.

In the case of notifications, the 2008 version made no mention of whether the customer or Cardinal sales would be informed of the threshold event, with the presumption being that both received notice.⁶²⁵ The 2009 version specifically provided that while sales will not receive a notification of the hold order, the customer will see on the invoice “held pending regulatory review.”⁶²⁶ The 2009 version does not preclude the sales team from receiving notice of held orders, only that the notice would not be system-generated. In the case of the customer, the notice of the hold would allow them to determine with some precision their thresholds, because they would know how much was dispensed when the held order was placed. Therefore, Cardinal was in effect giving customers the ability to circumvent the threshold control by manipulating order timing, requesting prospective threshold increases, etc. The subtlety with the Cardinal system was that Cardinal’s system undercut itself, thereby insulating the sales team from the appearance of directly subverting the threshold control.

The remainder of the procedure is devoted to detailing how sales personnel are to conduct “an inspection that looks for the Anti-diversion alert signals.”⁶²⁷ When conducting the inspection, the salesperson was to complete an online survey if the customer exhibited two or more of the alert signals to start the QRA Anti-Diversion team review process.⁶²⁸ If no signs of diversion were noted, then the sales representative completes the inspection and the form.⁶²⁹ However, the SOP is not explicit on when inspection and completion of the online survey is required. The SOP does contain an attachment with a memo to the Retail Independent Sales Professional

⁶²¹ *Id.* at § 4.1.

⁶²² *Id.* § 4.1 (Definition of “held order”).

⁶²³ *See* Sales Alert SOP 2008 at § 4.1 (Definition of “held order”).

⁶²⁴ Sales Alert SOP 2009, § 4.1 (Definition of “held order”).

⁶²⁵ *See* Sales Alert SOP 2008 at § 4.1 (Definition of “held order”).

⁶²⁶ Sales Alert SOP 2009 at § 4.1 (Definition of “held order”).

⁶²⁷ Sales Alert SOP 2008 at § 4.2 (Procedures for Reporting).

⁶²⁸ Sales Alert SOP 2008 at § 4.2(2) and (3).

⁶²⁹ *Id.*

describing “*The Highlight Report*,” which is the unofficial name.⁶³⁰ It shows the current monthly purchases of controlled substances as compared to a three-month rolling average of controlled substances sales. Customers are then flagged based upon the percentage increase in sales. There are three tiers: Watch List (5% or at least a \$2,500 increase), Yellow Flag (10% or at least a \$5,000 increase), and Red Flag (15% or at least a \$10,000 increase). Only Red Flag customers required an immediate store visit. Any customer that falls in the Watch List or Yellow Flag categories for three consecutive months was escalated to Red Flag status.

On a first pass, the Alert Signals inspection process looks adequate on paper. A closer inspection, however, reveals that in actuality, the process is filled with “loopholes” that allowed Cardinal to claim the company engaged in close customer monitoring without doing it. This is supported by Chris Lancot’s, Vice President of Sales for the Central Region, stated belief that sales did not have a role in ensuring compliance with the requirements the CSA, but simply complied with the directions provided by the QRA department.⁶³¹

First, the primary difference between the 2008 and 2009 versions can be seen in the list of “Anti-diversion alert signals” with the 2009 version being much more expansive and covering ambulatory surgical centers, and physicians’ offices, as well as pharmacies.⁶³² Conspicuously missing, however, from either version are pain management and other clinics; areas where significant diversionary activity already was occurring.

Second, regardless of its stated purpose, the Highlight Report process was a notification mechanism, alerting sales representatives to which customers were attracting attention through their purchasing patterns of controlled substances. The Highlight Report process did not require any action until the customers hit Red Flag status. Therefore, the astute sales representative could work with customers to either increase the thresholds or rework the ordering patterns such that the flag did not trip. Furthermore, Watch List or Yellow Flag customers only reached Red Flag status after three consecutive months. By adjusting ordering patterns, a customer could permanently avoid being on either list by simply adjusting hitting the Watch List or Yellow Flag for two months in a row with an intervening “normal” third month, which would reset the three-month trigger again.

Third, the Highlight Report compares the current month to a three-month rolling average of controlled substances sales, but it is unclear what makes up “controlled substances sales.” If Cardinal included all controlled substances sales across all customers, including chain stores, then the average would be so high as to render it meaningless for smaller independent retail pharmacies which appear to be the target of this process.

Fourth, despite the 2009 change from being directed to all Sales Professionals versus Retail Independent Sales Professionals in the Highlight Report memorandum, the system still does not appear to apply to chain store pharmacies.⁶³³ If it did, the sales increase limits are set so low that large chain customers such as Giant Eagle, CVS and Walgreens would need site visits on a monthly basis. Therefore, it appears that the change in addresses was done to avoid potentially difficult questions from a regulatory inspector or external auditors, about the limited application of the program. It is an important distinction that the Special Committee Report

⁶³⁰ See Sales Alert SOP 2008 at Attachment 1 (A memo dated 12/05/2008 from Tom DeGemmis, SVP to the Retail Independent Sales Professionals). The 2009 version contains virtually the same memo but addressed to “Sales Professionals” and dated 5/11/2009. See Sales Alert SOP 2009 at Attachment 1.

⁶³¹ See S. Lancot Deposition at 46:5-18 (Oct. 10, 2018).

⁶³² Sales Alert SOP 2009 at § 4.2.1 (Alert signals).

⁶³³ Sales Alert SOP 2008 at Attachment 1; *but cf.* Sales Alert SOP 2009 at Attachment 1.

seems to gloss over.⁶³⁴ In toto, as a diversion prevention control, the Alert Signal process is so easily circumvented that it cannot be considered much of a control at all.

E. On-site Investigations

Investigating potentially suspicious orders is a key component of any SOM program. Cardinal, however, did not establish a formal documented process for investigations until 2008.⁶³⁵ The SOP was intended to “provide guidance to CAH employees by outlining the steps involved in the conduct of on-site investigations of Cardinal Health's customers to obtain information regarding their potential risk for diversion of regulated drugs.”⁶³⁶ This process begins when “[a] pharmacist evaluates each threshold event according to established procedures and documents a request for an onsite investigation by changing the status of the event in the SOM system to ‘Site Visit.’”⁶³⁷ Presumably, this is a reference to the Detecting and Reporting Suspicious Orders and Responding to Threshold Events procedure, but the Investigations SOP fails to make a clear linkage.

The SOP also notes that the “Inspectors have no authority to require compliance with any request” for information.⁶³⁸ Given that Cardinal entered into purchasing contracts with customers, a normal compliance provision involves the inclusion of a “right to audit clause.” These clauses give a party the right to obtain documentation and conduct interviews. Without such authority, on-site investigators faced a substantial hurdle to obtaining the necessary information.

Finally, the investigative resolution process is not clear. Once the report was completed and the Director rendered a final recommendation, the next step was:

- A decision to continue the sale of regulated drugs to the customer requires an evaluation of the customer's threshold limits for regulated drugs and adjustments when supported by findings documented in the case. The Director, or another pharmacist, shall conduct such an evaluation and adjust thresholds appropriately.
- A decision to discontinue the sale of regulated drugs to the customer requires the termination of the customer from the Cardinal Health system and notification to state and federal regulatory bodies.⁶³⁹

What is not clear is how the decisions are implemented. For example, the Director clearly is not the person managing the termination, but the SOP is silent on who outside of QRA receives notice. Thus, as written the SOP is a dead-end process that results in the creation of a file but no apparent real action.

⁶³⁴ See Special Committee Report at 13.

⁶³⁵ See Cardinal Health, On-site Investigations, PDQRA-CAD-C008 (Nov. 5, 2009) (Initial issue of the procedure was 12/22/2008) CAH_MDL_PRIORPROD_DEA12_00014535. [“On-site Investigations SOP’s”].

⁶³⁶ *Id.* at § 1.1.

⁶³⁷ *Id.* at § 4.2.1.

⁶³⁸ *Id.* at § 4.3.5.

⁶³⁹ *Id.* at § 4.4.3.

F. QRA SOM Customer Analytics General Work Instructions

In January 2013, Cardinal developed a set of general work instructions covering SOM Customer Analytics.⁶⁴⁰ Typically, Work Instructions are a type control document issued as a supplement to the more formal policies and procedures and are intended to provide additional guidance to employees on how to implement specific policies and procedures.⁶⁴¹ As Todd Cameron noted, this particular set of Work Instructions provided “action-oriented details” in addition to what was set out in Cardinal’s existing set of SOM policies and procedures.⁶⁴² However, the General Work Instructions are not referenced in Cardinal’s standard operating procedure on detecting and reporting suspicious orders.⁶⁴³

These General Work Instructions outlined “the assessment and adjustment process” from beginning to end.⁶⁴⁴ As described, “[t]he initiation of an assessment could result from an early dialogue notice, held order, or proactive communication from the customer or sales department. The assessment could conclude with no change of a threshold limit, an increase or decrease of a threshold limit, resolution of a held order, and/or the report of a suspicious order to DEA.”⁶⁴⁵

The General Work Instructions also outline “the sequence of steps and corresponding decisions that should generally occur for each type of assessment.”⁶⁴⁶ These steps included:

- Determining the assessment was warranted;⁶⁴⁷
- Assessing a customer’s objective criteria;⁶⁴⁸
- Conducting an empirical review to “assess the reasonableness of the information and underlying basis for the threshold limit increase; and”⁶⁴⁹
- Determining eligibility for the increase based on the objective criteria and empirical review.⁶⁵⁰

⁶⁴⁰ See generally Cardinal Health, *QRA SOM Customer Analytics General Work Instructions*, (Sept. 20, 2013) (This is a revised version of the work instructions originally effective on January 15, 2013), CAH_MDL2804_00012249 [“SOM Analytics WI”].

⁶⁴¹ Work Instructions are a format that typically is employed in a manufacturing setting. In the manufacturing setting, Work Instructions usually are developed, revised and approved using the standard document control processes applicable to policies and procedures. See generally Appendix B, Figure 1 *infra*. However, here it seems that Cardinal considered the General Work Instructions to be “working guidelines.” See Email from K. Howenstein to K. Anna-Soisson, FW: assistance, (Apr. 28, 2014), CAH_MDL2804_00012244.

⁶⁴² See State of Montana, Office of the Attorney General, Office of Consumer Protection Deposition of Todd Cameron, 118:2-119:5, (Sept. 26, 2018).

⁶⁴³ See Cardinal Health Standard Operating Procedure Pharmaceutical Distribution, Detecting and Reporting Suspicious Orders and Responding to Threshold Events, PDQ-CAD-C007 (Jul. 18, 2014), CAH_MDL2804_000122245. For example, the General Work Instructions are not listed in the SOP’s references and related documents section although another SOP (PDQRA-CAD-C008) is listed.

⁶⁴⁴ See SOM Analytics WI at 1.

⁶⁴⁵ *Id.*

⁶⁴⁶ *Id.*

⁶⁴⁷ See *id.* at 2-4

⁶⁴⁸ See *id.* at 5-6.

⁶⁴⁹ *Id.* at 6.

Although the objective and empirical criteria are detailed and appear robust on their face, the Work Instructions provided a substantial “loophole” that allowed Cardinal to ignore situations where “the customer fails the objective criteria, and after empirical review, there is no justification for the threshold adjust[ment].”⁶⁵¹

According to the Work Instructions, [REDACTED]

[REDACTED]⁶⁵⁴ Therefore, despite making modifications to the SOM program after its 2012 DEA settlement, Cardinal persisted in maintaining practices that served to undercut those modified controls and gave the company the ability to apply its anti-diversion controls on a discretionary basis as demonstrated by these General Work Instructions.

In April 2014, Ullrich Mayeski, Cardinal’s Director of Investigations in Quality and Regulatory Affairs, asked Kim Howenstein and Kimberly Anna-Soisson to develop a presentation for Mr. Mayeski to give to Cardinal’s Compliance Officers at the distribution centers that they could “share with the DEA on-site and walk through with them during the inspection” to address “basic [SOM] components, particularly what happens when a threshold hits and cutting/reporting.”⁶⁵⁵ However, the June 2018 version of that presentation does not mention the ability to [REDACTED] found in the Working Instructions, except in the speaker’s note section.⁶⁵⁶

In an email to Ms. Howenstein, Ms. Anna-Soisson, Regulatory Management Manager for Cardinal, stated she “intentionally left out the part about [REDACTED],” because she “did not want to draw attention to the practice but agree that the CO’s should know it exists.”⁶⁵⁷ Ms. Anna-Soisson went on to add that she wondered “if we should take out any reference to the working guidelines since we don’t produce those.”⁶⁵⁸

⁶⁵⁰ *Id.* at 9.

⁶⁵¹ *See id.* at 5, § 4.a.c.i.

⁶⁵² *See id.* 7, § 4.a.c.i.vi.

⁶⁵³ *See id.* at 15 (Appendix 5, Table 5: Customer Release Percentage); *see also* Appendix D, Figure 2 *infra* (showing the actual table).

⁶⁵⁴ *See id.* 7, § 4.a.c.i.vi.

⁶⁵⁵ *See* Email from U. Mayeski to K. Anna-Soisson and K. Howenstein, assistance, (Apr. 28, 2014), CAH_MDL2804_00012244.

⁶⁵⁶ *See* Cardinal Health Quality & Regulatory, Cardinal Health Suspicious Order Monitor (SOM): Program Overview, 6 (Jun. 2018), CAH_MDL2804_00012954.

⁶⁵⁷ *See* Email from K. Howenstein to K. Anna-Soisson, RE: Adjustments made can you take a quick look, (May 1, 2014) (Ms. Howenstein’s email contains a copy of the text of Ms. Anna-Soisson’s original message.), CAH_MDL2804_00012953.

⁶⁵⁸ *Id.*

Since the General Work Instructions are written standards governing how Cardinal's SOM program operated and by the documents own words, applied to "all individuals who have the ability and/or direct responsibility for assessing and adjusting customer threshold limits,"⁶⁵⁹ it is concerning that Cardinal did not specifically reference them in the SOM SOPs and apparently had a policy not to disclose them. Regardless of the intent behind these actions, Cardinal, by making no reference to these General Work Instructions was maintaining a *sub rosa* process that failed to meet the standards of an effective compliance program.

10.6 Accountability - Consistent Enforcement

10.6.1 Cardinal does not enforce the standards of the program, and thus there is no real accountability for the program's lack of effectiveness.

While Cardinal "removed" certain staff members from positions of authority for controlled substances compliances, these staff members were simply transferred to other parts of the organization and thus never held accountable for their roles in the program's failure:

1. **Stephen Reardon** – Mr. Reardon, who oversaw the SOM program prior to 2007 was removed from controlled substances compliance responsibilities in 2007 when Michael Moné took over. However, he was shifted into a Vice President of Regulatory Operations where he remained until leaving Cardinal in 2016.
2. **Michael Moné** - Mr. Moné was involved with the anti-diversion team until 2012. He was replaced because "evaluation[s] of customers and orders had been heavily focused on the clinical expertise and subjective judgment of the pharmacists in the anti-diversion group."⁶⁶⁰ However, he was transferred to Cardinal's law department where he continues to service Cardinal as Vice President and Associate General Counsel.⁶⁶¹
3. **Steve Morse** – Mr. Morse was the Director of Supply Chain Integrity until he was removed in 2012 to become Quality and Regulatory Manager under Stephen Reardon in Regulatory Operations. He was transferred because he was "not as strategic as his former position required and there were questions about his judgment."⁶⁶² However, his new role requires "good judgment" as a requirement of the job.
4. **Gilberto Quintero** – Mr. Quintero was Senior Vice President of Quality and Regulatory Affairs from 2009 to 2015.⁶⁶³ In 2015, he was promoted to Chief Quality and Regulatory Affairs Officer – Pharmaceuticals & Medical Devices.⁶⁶⁴

Cardinal hired Craig Morford in 2008 to be its Chief Compliance Officer "with a mandate to establish a premier anti-diversion system."⁶⁶⁵ Mr. Morford remains Cardinal's CCO, as well as being its Chief Legal Officer;

⁶⁵⁹ See SOM Analytics WI at 1.

⁶⁶⁰ See Special Committee Report at 34-35.

⁶⁶¹ See Organization Chart 2012-2015, P1.4592.

⁶⁶² See Special Committee Report at 34-35.

⁶⁶³ *Id.*

⁶⁶⁴ See Gilberto Quintero LinkedIn Profile, <https://www.linkedin.com/in/gilbertoquintero/> (last accessed Jan. 29, 2019).

however despite having failed in his mandate to establish the premier anti-diversion and being the senior most officer responsible for controlled substances compliance, Cardinal has failed to hold him accountable.

By failing to hold these individuals accountable for the controlled substances program's established lack of effectiveness, Cardinal's compliance program is merely words on paper that do not meet the statutory and regulatory requirements.

11 AmerisourceBergen Corporation

11.1 Background

AmerisourceBergen Corporation ("AmerisourceBergen" or "ABC") traces its origin back to 1871 and the founding of the Brunswick Drug Company.⁶⁶⁶ Through a series of mergers and acquisitions beginning with the Bergen Drug Company in 1969, the Amerisource Health Corporation in 2001, and finally H.D. Smith in January 2018, AmerisourceBergen has become a worldwide distributor with more than 20,000 employees.⁶⁶⁷ Ranked 12th on the Fortune 500 with revenues in excess of \$150 billion,⁶⁶⁸ ABC declares that:

We provide the pharmaceutical products and business solutions that improve access to care. We operate the backbone of the healthcare supply chain.⁶⁶⁹

ABC distributes controlled substances through two wholly owned subsidiaries, AmerisourceBergen Drug Corporation ("ABDC") and AmerisourceBergen Specialty Group ("ABSG"), with ABDC being dominant.⁶⁷⁰

⁶⁶⁵ See Special Committee Report at 9.

⁶⁶⁶ See AMERISOURCEBERGEN, *Our History* at <https://www.amerisourcebergen.com/abcnew/about-our-history> (last accessed Jan. 19, 2019).

⁶⁶⁷ *Id.*; see also AMERISOURCEBERGEN, *Who We Are* at <https://www.amerisourcebergen.com/abcnew/about-who-we-are> (last accessed Jan. 19, 2019); see also BUSINESSWIRE, *AmerisourceBergen Completes Acquisition of H.D. Smith*, (Jan. 3, 2018, 0800 EST) (At the time, H.D. Smith was the largest independent wholesaler in the U.S.) at <https://www.businesswire.com/news/home/20180103005161/en/AmerisourceBergen-Completes-Acquisition-H.-D.-Smith>.

⁶⁶⁸ FORTUNE 500, *Amerisource Bergen* (last accessed Jan. 21, 2019), <http://fortune.com/fortune500/amerisourcebergen/>.

⁶⁶⁹ See AMERISOURCEBERGEN, *Who We Are* at <https://www.amerisourcebergen.com/abcnew/about-who-we-are> (last accessed Jan. 19, 2019).

⁶⁷⁰ As used in this report, AmerisourceBergen or ABC includes two subsidiaries that distribute controlled substances. The first, AmerisourceBergen Drug Corporation or ABDC, "distributes pharmaceuticals products, equipment, and systems ... [and] serves healthcare providers, independent retailers, and pharmacies." See Company Overview of AmerisourceBergen Drug Corporation, BLOOMBERG at <https://www.bloomberg.com/research/stocks/private/snapshot.asp?privcapId=928736> (last accessed Jan. 21, 2019 3:50 PM ET). AmerisourceBergen Specialty Group "distributes medical products to healthcare providers ... [including] chemotherapy and supportive care products to oncology practices." See Company Overview of AmerisourceBergen Specialty Group, BLOOMBERG at <https://www.bloomberg.com/research/stocks/private/snapshot.asp?privcapId=27144580> (last accessed Jan. 25, 2019 9:12 AM ET).

Although prescription opioids account for less than 2% (\$3 billion) of ABC's total revenues,⁶⁷¹ this translates into a vast number of dosage units.

ABC's controlled substances program traces its origins back to Bergen Brunswig's system. At the formation of AmerisourceBergen in 2001, ABC adopted the Bergen Brunswig anti-diversion system.⁶⁷² During the review period, ABC, and its predecessor Bergen Brunswig have made periodic updates to its controlled substances program with the major milestones being 1997, 2007 and 2014.⁶⁷³

On the enforcement front, AmerisourceBergen Drug Corporation entered into a settlement with the DEA in June 2007 to resolve allegations that the company failed to maintain effective controls against diversion.⁶⁷⁴ As noted in ABC's press release announcing the settlement, "[t]he agreement requires the Company to implement an enhanced and more sophisticated order monitoring program in all AmerisourceBergen Drug Corporation distribution centers by June 30, 2007, after which the Company must pass several DEA inspections of the new program for the reinstatement to become effective [in August 2007]."⁶⁷⁵

Although the company has not been the subject of a second enforcement action by the DEA, ABC has undergone several subsequent DEA inspections that have noted deficiencies in ABC's recordkeeping processes.⁶⁷⁶ Furthermore, ABC has been embroiled in other serious compliance breaches, with the most notable being the 2018 anti-kickback settlement involving ABSG.⁶⁷⁷

⁶⁷¹ Written Statement of Steven H. Collis Chairman, President, and Chief Executive Officer AmerisourceBergen Corporation Before the Subcommittee on Oversight and Investigations Committee on Energy and Commerce U.S. House of Representatives at (May 8, 2018), <https://docs.house.gov/meetings/IF/IF02/20180508/108260/HHRG-115-IF02-Wstate-CollisS-20180508.pdf> ["Collis Statement"].

⁶⁷² See Email from C. Zimmerman to P. Ross, Emailing BOD CS TPs 8.10.17 (8.6.17) Final, 2 (Aug. 7, 2017), ABDCMDL00273269 -ABDCMDL00273270 ["BOD Talking Points"].

⁶⁷³ See BOD Talking Points at 2.

⁶⁷⁴ See Settlement and Release Agreement between the U.S. Department of Justice, Drug Enforcement Administration and AmerisourceBergen Drug Corporation (Jun. 22, 2007), ABDCMDL00279854 ["ABDC Settlement"].

⁶⁷⁵ Amerisource Bergen Corporate Press Release, *AmerisourceBergen Signs Agreement with DEA Leading to Reinstatement of Its Orlando Distribution Center's Suspended License to Distribute Controlled Substances* (Jun. 22, 2017), <http://investor.amerisourcebergen.com/news-releases/news-release-details/amerisourcebergen-signs-agreement-dea-leading-reinstatement-its>.

⁶⁷⁶ See Email from Steve Mays to D. May and C. Zimmerman, *FW: DEA Audit Tracking* (May 16, 2017), ABDCMDL00253868.

⁶⁷⁷ In September 2018, ABC and AmerisourceBergen Specialty Group ("ABSG") agreed to pay \$625 million and entered into a five-year Corporate Integrity Agreement ("CIA") to settle allegations of distributing misbranded oncology products and harvest[ing] "overfill" from the original vials of [chemotherapy] drugs ... [t]hat enabled the company to create more doses than it bought and generate at least \$99.6 million of extra profit. ... AmerisourceBergen was also accused of billing multiple doctors for individual vials, causing them to bill the government more than once, and paying kickbacks to induce doctors to buy drugs through the pre-filled syringe program." See Jonathan Stempel, *AmerisourceBergen to pay \$625 million in U.S. civil fraud settlement*, REUTERS (Oct. 1, 2018, 2:16 PM), <https://www.reuters.com/article/us-amerisourcebergen-settlement/amerisourcebergen-to-pay-625-million-in-u-s-civil-fraud-settlement-idUSKCN1MB3IT>.

11.2 Executive Summary

In contrast to both McKesson and Cardinal Health, AmerisourceBergen's culture is a paradox with the company presenting a different "public" and a "private" face. Understanding this paradox is central to understanding ABC's approach to controlled substances compliance.

Publicly, AmerisourceBergen maintains and apparently believes that ABC's controlled substances compliance program always was compliant. ABC further maintains that any changes made to the program, including those resulting from the 2007 settlement, were the result of ABC's cooperative working arrangement with the DEA, not a compliance breach.

Privately, ABC, and those responsible for the controlled substances program, not only were conversant with the DEA's expectations for distributors, but they worked to configure a program that only addressed the bare minimums and did not interfere with ABC's pursuit of ever-increasing revenues. Moreover, internally, ABC attempts to shift its responsibilities to maintain effective anti-diversion controls to the DEA, based on a lack of direct communication by the DEA to ABC. Thus, ABC's poor compliance culture substantially contributed to it ineffectually addressing its obligations as a distributor of controlled substances.

Expanding on that notion of dialog with the DEA, AmerisourceBergen developed the misguided narrative that it was entitled to regular communications with the DEA, including having the DEA supply it with information on diversionary customers and review its systems. Despite receiving contrary information directly from the DEA, this expectation persisted, and it simply ignores the reality that the responsibility to identify and report suspicious orders as well as maintain an effective anti-diversion program rests with the distributor and not the DEA.

Also, in contrast to both McKesson and Cardinal Health, AmerisourceBergen, beginning in 2007, applied and continues to apply effort and resources towards improving its controlled substances compliance program. Unfortunately, however, most of ABC's efforts dedicated to controlled substances compliance have been misapplied and thus ineffective at achieving a credible and workable anti-diversion program.

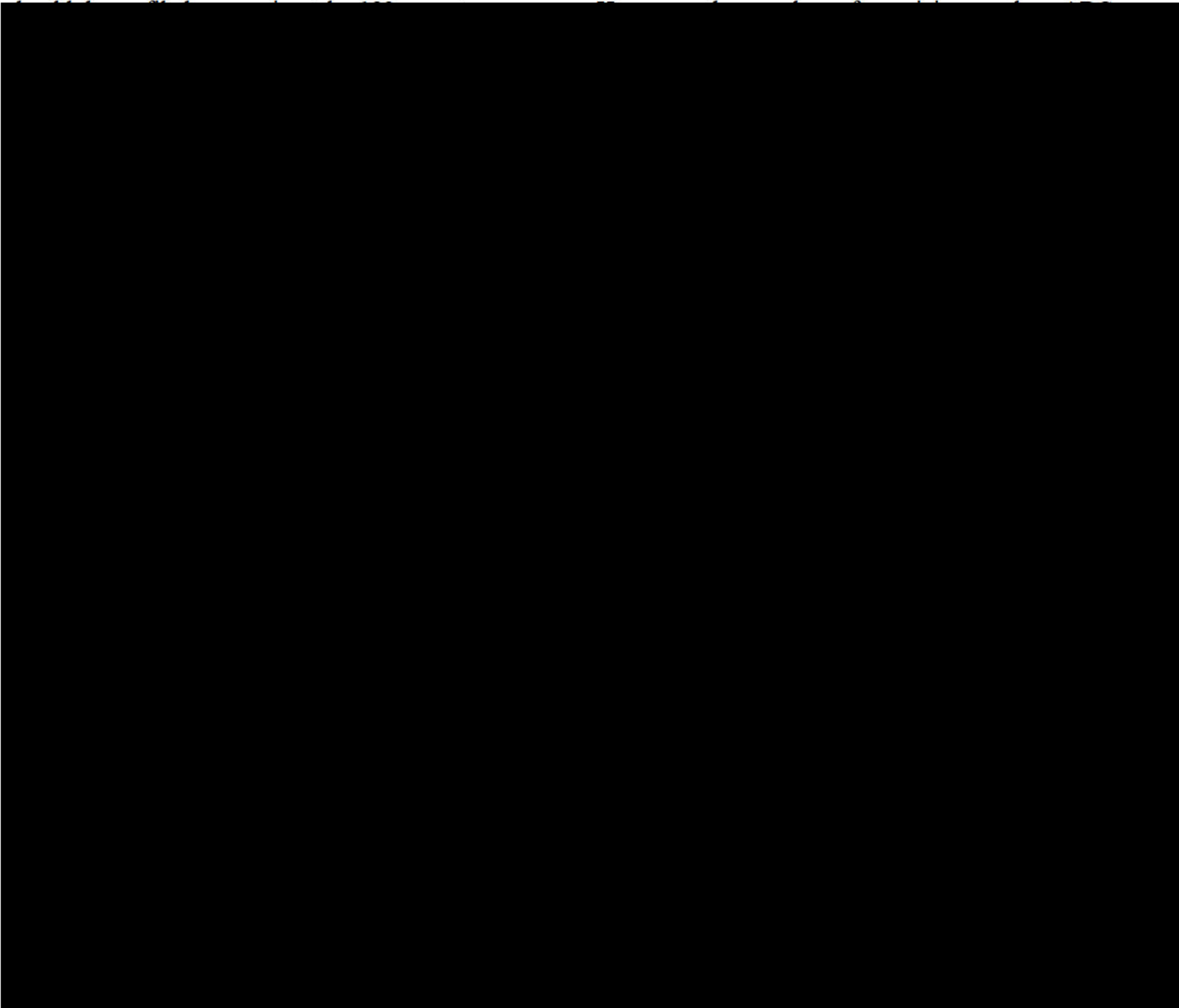
Poor program design and inconsistent application of the standards that were developed exacerbated the deficiencies in AmerisourceBergen's controlled substances program leading to a predictable outcome that ABC's program credibly failed to identify, report and stop suspicious orders.

11.3 Impact

An example of the failure to maintain a credible anti-diversion program can be seen in West Virginia. As the House Energy and Commerce Committee uncovered, between 2005 and 2016, AmerisourceBergen distributed 248.16 million dosage units of hydrocodone and oxycodone to West Virginia customers.⁶⁷⁸ [REDACTED]

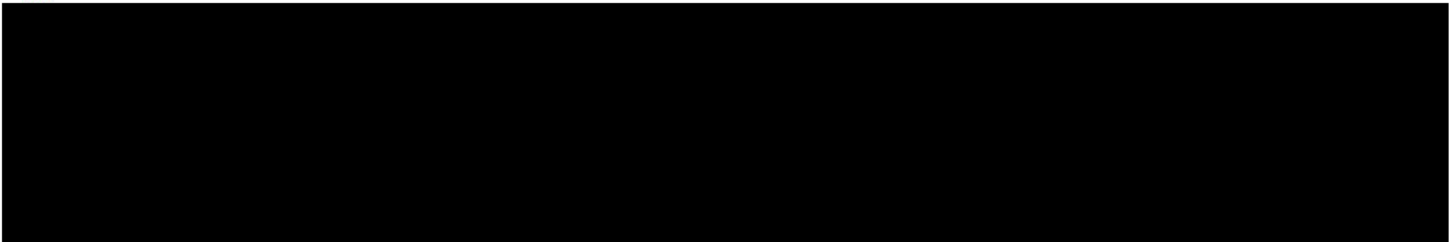
⁶⁷⁸ See U.S. House Energy & Commerce Committee Report, *Red Flags and Warning Signs Ignored: Opioid Distribution and Enforcement Concerns in West Virginia*, 115th Cong., 6 (Dec. 19, 2018), P1.2060 ["W.Va. Red Flags Report"].

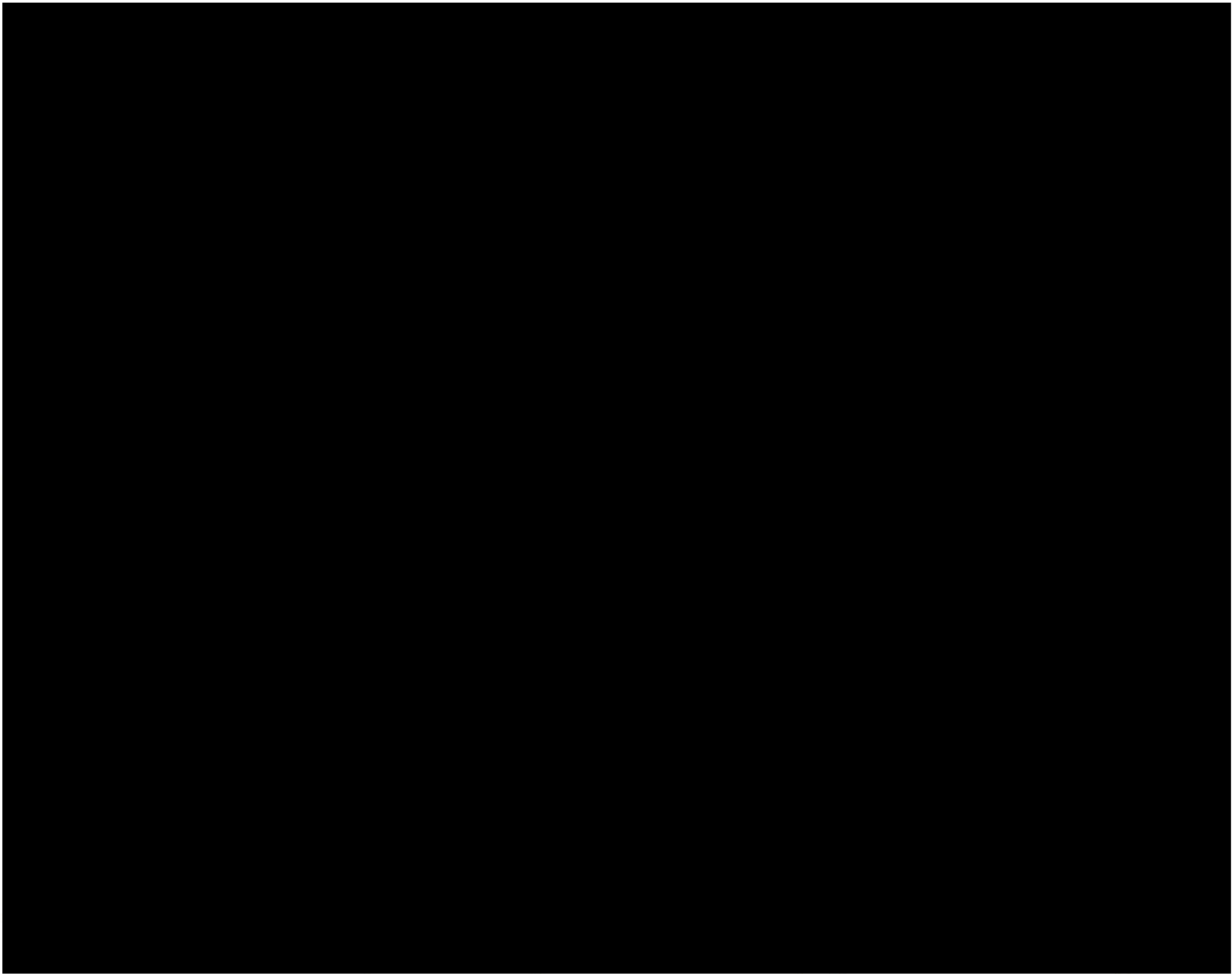
⁶⁷⁹ See Appendix E, Figure 3.



⁶⁸⁰ See Appendix E, Figure 4.

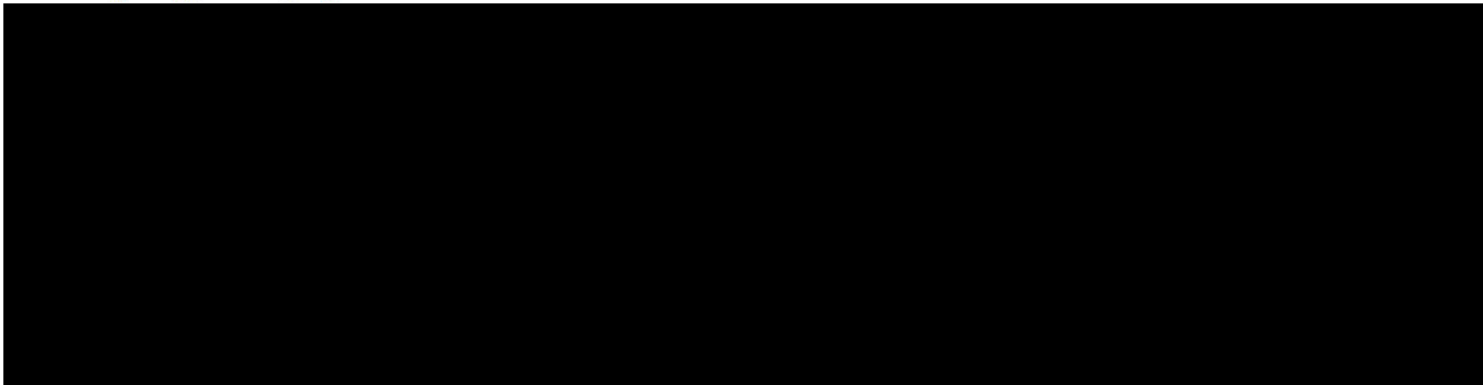
⁶⁸¹ *Id.* at Appendix E, Figure 4.





⁶⁸⁶ See *infra* Appendix E, Figure 1.

⁶⁸⁷ See *infra* Appendix E, Figure 1.



11.4 Company Commitment – Compliance Culture, Organization & Resources

11.4.1 The “private” versus “public” cultural paradox within AmerisourceBergen hampered its ability to create and maintain an effective controlled substances compliance program.

AmerisourceBergen’s culture throughout the period is characterized by a public versus private paradox. Thus, there are two faces to AmerisourceBergen: a “public” and a “private” face. Publicly, ABC acknowledges that distributors have a role in preventing diversion under the Controlled Substances Act. AmerisourceBergen also states publicly that it takes this role seriously and maintains that its controlled substances program is compliant even though it has changed and updated its program during the period. ABC continues to maintain that stance in the face of evidence about the program’s deficiencies documented and provided to it by outside parties (e.g., the DEA and FTI).⁶⁹⁷

In public, ABC’s current Chairman, President, and CEO, Steven Collis in hearings before House Energy and Commerce Committee’s Subcommittee on Oversight and Investigations, acknowledged that with respect to opioids, ABC’s “distribution role in the [drug supply] system is vital, yet limited, “ but that the company is “responsible for getting those medicines to tens of thousands of sites of care every day, including pharmacies, hospitals, and clinics, which administer or dispense the medicines on prescriptions written by licensed health

⁶⁹⁷ See, e.g., Email from Steve Mays to D. May and C. Zimmerman, FW: DEA Audit Tracking (May 16, 2017), ABDCMDL00253868; FTI Consulting, Inc., Health Solutions Practice, *AmerisourceBergen Corporation CSRA Process Review, Phase 1 Narrative Report*, 3 (Aug. 25, 2015), ABDCMDL00274105 [“FTI Narrative”]

care providers.”⁶⁹⁸ AmerisourceBergen contends that the company “worked with the DEA to enhance the system in 1998, and again in 2007, and have continually reviewed and improved it [the anti-diversion program], including a comprehensive 2015 revision to build on current data, respond to trends in prescription drug abuse, and adopt improved technological capabilities, including data-driven analytical tools.”⁶⁹⁹ This is the public face of AmerisourceBergen.

The public face of AmerisourceBergen also contends that its controlled substances compliance program always was compliant. For example, Chris Zimmerman, Senior Vice President Corporate Security and Regulatory Affairs and Chief Compliance Officer,⁷⁰⁰ in his deposition framed the DEA communications with the company, including during the 2007 settlement, as AmerisourceBergen simply was being asked to modify its program, as opposed to the fact that DEA was notifying ABC its program was operating contrary to DEA requirements.⁷⁰¹

On the private side, there was the expectation that ABC was entitled to get regular guidance from the DEA. For example, in his 2017 Board talking points, Mr. Zimmerman referenced the fact that ABC enhanced its controlled substances program “without participation or input from the DEA, as DEA had stopped communicating with industry at this point.”⁷⁰² However, the DEA was not obligated to participate in or provide input on ABC’s program, and doing so would have run contrary to the DEA’s longstanding position that it does not endorse particular systems or programs. Furthermore, Mr. Zimmerman’s talking points neglected the fact that ABC, not the DEA, is responsible for maintaining an effective program to prevent diversion.

This private face also can be seen in the “comprehensive 2015 revision” referenced in Mr. Collis’ Congressional testimony. In 2015, ABC engaged FTI Consulting, Inc.’s Health Solutions Practice (“FTI”) to:

review, map and document current state processes; identify any critical process gaps and areas for improvement; develop recommendations for process improvements and gap remediation; and scope the initial functional requirements for a technology solution(s) or technology enhancements identified as part of the recommendations.⁷⁰³

⁶⁹⁸ See Collis Statement at 3-4.

⁶⁹⁹ See *id.* at 7-8.

⁷⁰⁰ In October 2018, Mr. Zimmerman was replaced by Kathy Gaddes, who is Executive Vice President and Chief Compliance Officer for ABC. See AmerisourceBergen, Kathy H. Gaddes at <https://www.amerisourcebergen.com/abcnew/people/kathy-gaddes> (last accessed Jan. 22, 2019). Prior to that she was Executive Vice President and Chief Human Resources Officer. Mr. Zimmerman’s replacement coincides with ABSG’s September 2018 AKS settlement. See Discussion *infra*.

⁷⁰¹ See Chris Zimmerman Deposition, 139 (Aug. 3, 2018).

⁷⁰² See Board Talking Points at 2.

⁷⁰³ See FTI Consulting, Inc., Health Solutions Practice, *AmerisourceBergen Corporation CSRA Process Review, Phase 1 Narrative Report*, 3 (Aug. 25, 2015), ABDCMDL00274105 [“FTI Narrative”]

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[REDACTED]

This private face can also be seen in AmerisourceBergen's Code of Ethics and Business Conduct. The Code contains the normal company admonishments to employees:

- act ethically;
- comply with the law;
- protect the Company's tangible and intangible assets; and
- respect and ensure the safety of all Company employees.⁷⁰⁸

Despite its public acknowledgment by ABC's Chairman, President, and CEO, that distributors have a role in preventing diversion and the company takes this role seriously, AmerisourceBergen's Code does not prioritize its responsibilities regarding controlled substances and only briefly mentions controlled substances in passing noting:

Furthermore, each operating company may have additional policies and procedures that further clarify your ethical and legal obligations. For instance, associates of some business units of the Company are also required to comply with the Company's Marketing Code of Conduct. Other associates, **including all compliance-critical associates with ongoing authorization to access controlled substances**, and certain key management personnel, are subject to additional compliance training and annual screening.⁷⁰⁹

Instead of focusing on preventing controlled substances diversion, the Code spends proportionately more time on covering social media, company ownership of intellectual property, and compliance approval forms for gifts and donations. The Code also states that associates (employees) can be disciplined for not reporting suspected violations but does not impose an affirmative duty to report violations.⁷¹⁰ Thus, the Code itself illustrates this public/private cultural paradox in which ABC's actions surrounding controlled substances compliance are not aligned with the public perception it tries to maintain.

⁷⁰⁵ [REDACTED]

⁷⁰⁶ See *id.*

⁷⁰⁷ [REDACTED]

⁷⁰⁸ See AmerisourceBergen, *Code of Ethics and Business Conduct*, 3 (Mar. 2017), P-138 ["ABC COE"].

⁷⁰⁹ See ABC COE at 3 (emphasis added).

⁷¹⁰ See ABC COE at 5.

This goes beyond just the Code and permeates other ABC policies as well. For example, ABC states that “each associate has a responsibility by federal administrative law ... to report any diversion of any listed chemical or controlled substance from our company **by fellow associates.**”⁷¹¹ However, the Code does not require associates to report diversion or suspected diversion by **customers** as envisioned by the CSA. Therefore, the Code demonstrates that when ABC looks at diversion, it is more concerned with diversion by the company’s employees that directly affects ABC’s profitability, rather than diversion by its customers once the controlled substances are sold.

Consequently, ABC’s company culture and its “private face,” have hampered ABC’s ability to reflect on the effectiveness of its program objectively. They also have created a contradictory paradigm in which ABC, on the one hand, maintains its program is fully compliant with the DEA’s requirements, and yet, on the other hand, the company makes periodic changes to the CSRA organization and the program’s core. Consequently, ABC’s attempt “to walk in two worlds” simply has hamstrung its ability to develop and implement an effective program to detect and report suspicious orders and to potentially alleviate the diversionary activities undertaken by some of its customers.

11.4.2 AmerisourceBergen has failed to optimize its organizational connections between ABC’s controlled substances program and ABC’s Corporate Compliance program.

Since its inception in 2001, AmerisourceBergen has had three Chief Compliance Officers. The three are:

- Debbie Schwartz, Associate General Counsel & Chief Compliance Officer (2001 to 2012);⁷¹²
- Christopher Zimmerman, Senior Vice President Corporate Security and Regulatory Affairs & Chief Compliance Officer (2012 to 2018);⁷¹³ and
- Kathy Gaddes, Executive Vice President and Chief Compliance Officer (2018 to present).⁷¹⁴

Of the three, it is Mr. Zimmerman’s tenure as Chief Compliance Office beginning in 2012 that has had the greatest detrimental effect on the company’s controlled substances and corporate compliance programs. With the expansion of Mr. Zimmerman’s role to head up both the controlled substances and corporate compliance programs, AmerisourceBergen had an opportunity to dramatically raise the profile and importance of controlled substances compliance within the organization. Unfortunately, Mr. Zimmerman and ABC squandered that opportunity.

Despite having regular contact with ABC’s Board of Director’s Audit Committee, Mr. Zimmerman failed to use those interactions to highlight and address controlled substances compliance. As he testified in his deposition,

⁷¹¹ See AmerisourceBergen, *Associate Responsibility to Report Diversion*, S&RC 12.01 (May 13, 2016) (emphasis added) (The policy was first made effective on 10/1/2005), ABDCMDL00156065.

⁷¹² See D. Schwartz LinkedIn Profile, <https://www.linkedin.com/in/debbieswartz/> (last accessed Jan. 26, 2019).

⁷¹³ See C. Zimmerman Deposition at 20:18-20; see also AmerisourceBergen, Kathy H. Gaddes at <https://www.amerisourcebergen.com/abcnew/people/kathy-gaddes> (last accessed Jan. 22, 2019).

⁷¹⁴ See AmerisourceBergen, Kathy H. Gaddes at <https://www.amerisourcebergen.com/abcnew/people/kathy-gaddes> (last accessed Jan. 22, 2019).

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In addition, despite Mr. Zimmerman also being ABC's Chief Compliance Officer, evidence suggests that CSRA, in general, lacked credibility and authority within the AmerisourceBergen organization. FTI highlighted this in its 2015 CSRA review noting

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FTI's report is not the only evidence of the weakness of CSRA. It also can be seen during the development of CSRA talking points for the Regional Vice Presidents outlining ABC's new threshold approach in 2009. The new threshold approach involved factoring both customer size and controlled substances purchasing ratios.

Describing the new approach, Mr. Zimmerman wrote that "[i]f this [approach] is acceptable, we will undertake a review of the top 25 purchasers of HY and OX and adjust thresholds accordingly"717 Although the CSRA and Mr. Zimmerman had full responsibility for the controlled substances compliance program, this statement shows that Mr. Zimmerman felt he needed approval to act. This is a sign that either Mr. Zimmerman did not understand his role as Chief Compliance Officer or that he knew CSRA's actual authority was extremely limited.

A good compliance officer normally does not seek approval to act within the department's delegated authority, especially when the action can be easily modified or reversed (e.g., no lasting impact). Therefore, I would have expected a statement that he was implementing the new threshold approach after engaging in appropriate consultation with the business, rather than his statement here.

11.4.3 While the number of personnel assigned to AmerisourceBergen's CSRA function grew over time, the company did not utilize those resources to its best advantage.

From a total paper headcount perspective, AmerisourceBergen has a sizable number of resources in its Corporate Security and Regulatory Affairs ("CSRA") function when compared to its peers (e.g., Cardinal and McKesson). However, the numbers alone do not tell the whole story. First CSRA has more duties than just handling controlled substances compliance and second, CSRA's organizational structure is so convoluted that the division of responsibilities within the CSRA team is unclear. The result is that the organizational structure of CSRA is a net negative that contributes to program inefficiency and inconsistency.

⁷¹⁵ See C. Zimmerman Deposition at 252:4-253:6.

⁷¹⁶ See FTI Narrative at 8 (emphasis added).

⁷¹⁷ See Memorandum from C. Zimmerman to E. Hazewski, *et al.*, *RVP Talking Points*, 2 (Jan. 19, 2009) (emphasis added), ABDCMDL00000169 ["RVP Talking Pts. 2009"].

Within ABC, the CSRA department has an expansive role that includes diversion control, regulatory compliance and auditing, investigations, complaints, business continuity and physical security.⁷¹⁸ In May 2012 those duties expanded once more when Mr. Zimmerman become ABC's, Chief Compliance Officer.

In the areas of diversion control, the CSRA Diversion Control Group was first established in 2007.⁷¹⁹ From 2008 to 2014, the group was headed up by Edward Hazewski, Director of Diversion Control and Security.⁷²⁰ Although at the outset the Diversion Control Group reported directly to Mr. Zimmerman as the head of CSRA, its reporting relationship has devolved several times over the years, including reporting for a time to the Drug Distribution Group, thereby reducing its visibility within CSRA. Nor did the changes translate into significant headcount increases. The group always has been small ranging in size from a minimum of 3 in 2009⁷²¹ to a maximum of 17 in 2017.⁷²²

In stark contrast, CSRA's Drug Distribution Group, headed by Stephen Mays, ranged in size from a minimum of ~32 in 2009⁷²³ to a maximum of ~43 in 2017.⁷²⁴ As recounted by Mr. Mays, from 2007 to 2015, the Drug Distribution Group (currently known as the Pharmaceutical Distribution and Global Sourcing Group) had primary responsibility for the Order Monitor Program or OMP.⁷²⁵ Therefore, since 2007 the Drug Distribution Group has played a significant role in ABC's controlled substances compliance program.

Even when most of ABC's diversion control efforts were centralized under the auspices of David May, Vice President of Security and Diversion Control starting in 2014 and completed 2015, ABC failed to clarify roles and responsibilities sufficiently. Mr. May, together with his seven direct reports in 2015:

Directs overall Diversion Control Program for ABC and all subsidiaries. Initiates and manages program initiatives to ensure that the drug company and the other business units that distribute controlled substances and listed chemicals are operating within established law and regulation. Tracks and responds to changes in prescription drug abuse trends and regulatory requirements. Assists legal in preparing and responding to official state and federal requests and subpoenas for information and documentation. Provides information and training to customers relative to diversion control best practices. Evaluates customer due diligence investigations and take appropriate actions to prevent illegal diversion. Develops and manages anti-diversion education and training programs tailored to company business units and functions. Maintains working relationships with drug manufacturers and other industry stakeholders.⁷²⁶

⁷¹⁸ See FTI Narrative at 6.

⁷¹⁹ See BOD Talking Points at 2.

⁷²⁰ See Edward Hazewski Deposition, 21:4-12 (Oct. 25, 2018).

⁷²¹ See CSRA Organizational Chart, Policy 1.1 (Oct. 1, 2009), ABDCMDL00364269.

⁷²² See CSRA Organization Chart, ABDCMDL00017114 (Oct. 2017).

⁷²³ See CSRA Organizational Chart, Policy 1.1 (Oct. 1, 2009), ABDCMDL00364269.

⁷²⁴ See CSRA Organization Chart, ABDCMDL00017114 (Oct. 2017).

⁷²⁵ See Stephen Mays Deposition, 66:21-67:7 and 200:24-202:23 (Oct. 24, 2018) (Stephen Mays was the top diversion control person in 2007).

⁷²⁶ See CSRA, Diversion Program Assignments, 1 (May 2015), ABDCMDL00247169; see also Senior Director, CSRA

Even during the centralization period (2014-2015), when CSRA began outlining the new Diversion Control and Federal Investigations role, it was clear that Pharmaceutical Distribution would continue to play a prominent, if unclear, role in ABC's diversion prevention program.⁷²⁷ For example, as the responsibility for customer communications illustrates, this responsibility was vested jointly between Diversion Control and Pharmaceutical Distribution, but with no clear delineation of roles and responsibilities.⁷²⁸ As FTI noted in its report:

The CSRA resources that we interacted with were very knowledgeable with respect to their subject areas, although at times it was challenging to nail down their specific scope of responsibilities. Because **the roles and responsibilities of CSRA personnel are somewhat ill-defined**, CSRA team members sometimes end up performing activities outside of their purview which **detracts from the utilization of these resources**.⁷²⁹

Further complicating matters was the reliance by CSRA on the ABC sales team, which in addition to interacting directly with customers and growing the company's "top line", also was responsible for collecting the Form 590s, a major part of ABC's due diligence efforts⁷³⁰, as well as reporting "red flags" of diversion back to CSRA, but these "extra duties" were not part of the overall sales representatives' compensation plan.⁷³¹ According to Nathan Elkins, like most sales teams, ABC's representative compensation was tied to meeting quotas (e.g., sales targets).⁷³² However, up until FY 2017, the sales of controlled substances, including opioids, were included in the sales representatives' compensation plan.⁷³³ It was only after 2017 that opioids were "carved out" of the mix.⁷³⁴ Therefore, before the FY 2017 "carve out," greater opioid sales could lead to greater compensation for ABC sales representatives.

The use of sales targets and quotas by a distributor or pharmaceutical company is not *per se* wrong. However, in the case of ABC, such a heavy reliance on the sales team to meet CSRA's compliance needs without providing a corresponding compliance performance objective disincentivized ABC's sales team from reporting compliance issues. It follows the well-worn corporate maxim that what gets measured and rewarded gets done.

Diversion Control and Federal Investigations, 2014 Performance Work Plan (Apr. 29, 2014), ABDCMDL00158306 and ABDCMDL00158307.

⁷²⁷ See, e.g., Email from C. Zimmerman to E. Hazewski and S. May., FW: Updated (Feb. 12, 2014) ("we decided ... [to] get [in] a room and come to a consensus of who is going to do what."), ABDCMDL00291410; CSRA Task Survey, ABDCMDL00246985; E. Hazewski, *Diversion Control Program Roles* (Feb. 10, 2014), ABDCMDL00291411.

⁷²⁸ See E. Hazewski, *Diversion Control Program Roles* (Feb. 10, 2014), ABDCMDL00291411; FTI Findings at 20.

⁷²⁹ See FTI Narrative at 7-8 (emphasis added).

⁷³⁰ For a description of and discussion about the CSRA Form 590s see *infra* Section 10.5.4.

⁷³¹ See Nathan Elkins Deposition, 225-230 (Nov. 14, 2018); see *id.* at 152:6-15.

⁷³² See *id.* at 135:14 to 136:16. According to his LinkedIn profile, Mr. Elkins, currently an ABC Sales District Director, was a Retail Account Manager from 2005 to 2011. See Nathan Elkins LinkedIn Profile, <https://www.linkedin.com/in/nathan-elkins-8a1b0718/> (last accessed Mar. 16, 2019).

⁷³³ See *id.* at 142:23-143:7.

⁷³⁴ See *id.* at 143:1-7 and 144:4-9.

Finally, poor utilization and the lack of clear responsibilities ultimately increased workload and reduced the program's overall effectiveness.

11.5 Program Core – Requirements, Education, Detection & Corrections

11.5.1 Prior to 2007, AmerisourceBergen's two-part controlled substances program was at best rudimentary, and not compliant with DEA regulatory requirements.

Like other distributors, ABC's pre-2007 controlled substances program had an extremely limited focus. The program's primary objective was to notify the DEA about suspicious orders rather than to create a holistic anti-diversion program. According to Mr. Zimmerman, in 1997, Bergen Brunswig, ABC's predecessor, revamped its order monitoring program ("OMP") "to account for the individual pharmacies purchasing levels by volume."⁷³⁵ Prior to that date, both large and small pharmacies were lumped together into a single peer group.⁷³⁶

During that period (1997 to 2007), ABC did not hold or investigate potentially suspicious orders (or as ABC referred to them - orders of interest), as the practice was to ship the orders at night, and then the next day any orders that were identified as suspicious were reported to the DEA.⁷³⁷ Although the process was a two-step process ("It was an excessive order report that was produced monthly to send to DEA, and then we also had a manual process at the distribution centers where the order fillers would identify suspicious orders and report those.") the clear emphasis was on processing orders as quickly as possible to meet customer demands.⁷³⁸

A. Excessive Order Reports

As the primary SOM control pre-2007, excessive order reports were generated when an ABC customer exceeded its threshold or monthly allocation for a controlled substance. According to Stephen Mays, Senior Director, Pharmaceutical Distribution and Global Sourcing, an "excessive order" was "an order that would have exceeded those parameters that were built into the system to produce those reports," while a "suspicious order" was "[a]nything that met those guidelines and the regulation that could be a suspicious order."⁷³⁹ Therefore, according to Mr. Mays, not every excessive order was a suspicious order.

Underpinning the excessive order reports were product thresholds. According to Mr. Zimmerman, ABC has always employed some form of thresholds.⁷⁴⁰ Prior to 1998, ABC simply generated an average volume per

⁷³⁵ See BOD Talking Points at 2.

⁷³⁶ See BOD Talking Points at 2.

⁷³⁷ See Chris Zimmerman Deposition at 110:16-22.

⁷³⁸ See C. Zimmerman Deposition at 108:19-109:4.

⁷³⁹ See Steve Mays Deposition, 131:5-16 (Oct. 24, 2018).

⁷⁴⁰ See C. Zimmerman Deposition at 111:12-16.

month per drug category and then added the multiplier of three.⁷⁴¹ Orders above that level would be on the excessive order report.⁷⁴²

Beginning in 1998 through 2007, ABC refined its threshold calculations even further, but still retained the multiplier of three. According to Mr. Zimmerman, ABC examined a pharmacy's purchasing history and calculated a rolling four-month average that was then tripled to create the suspicious order threshold.⁷⁴³

While this new calculation improved upon the earlier version by reducing the base threshold for smaller pharmacies, the arbitrary tripling of the base threshold for all pharmacies could still allow pharmacies to order excessively large quantities of opioids and other controlled substances without being flagged. Having a more refined "buffer" built into the threshold would have reduced that risk. Given the fact that ABC was not holding, but merely reporting, any excessive orders, having a more refined threshold likely would have achieved nothing during the time period before 2007. However, the lack of a refined threshold together with ABC's failure to hold and investigate excessive orders resulted in a deficient SOM and diversion prevention program during this period.

B. Manual Distribution Center Process

The manual distribution center process (the second step) cannot remedy the deficiencies in ABC's pre-2007 program. Like Cardinal Health, ABC relied on "pickers" to identify potentially suspicious orders. ABC asked its "distribution center employees that work in the cages and vaults if they see an unusually large order or frequency or pattern that they feel could be potentially suspicious, then they are to report that."⁷⁴⁴

However, ABC's policy outlining an associate's responsibility for diversion further undermines reliance on the cage and vault employees as an effective control. According to Policy S&RC 12.01 which initially became effective in October 2005, "each associate has a responsibility by federal administrative law ... to report any diversion of any listed chemical or controlled substance from our company **by fellow associates**."⁷⁴⁵ This policy does not require associates to report customer orders of unusual size, frequency or pattern. Therefore, at the bare minimum, this policy incompletely covers an employee's affirmative duties pertaining to controlled substances compliance.

11.5.2 AmerisourceBergen's Order Monitoring Program ("OMP") between 2007 and 2016 was rendered ineffective by a combination of poor design and inconsistent application.

In 2007, AmerisourceBergen proceeded to modify its controlled substances compliance program. Mr. Zimmerman described the changes as:

⁷⁴¹ *Id.* at 121:12-20.

⁷⁴² *Id.* at 121:18-20.

⁷⁴³ *Id.* at 122:13-23.

⁷⁴⁴ *Id.* at 114:22-115:6.

⁷⁴⁵ *See Associate Responsibility to Report Diversion*, S&RC 12.01 (emphasis added).

DEA wanted us to include a more in-depth due diligence process in addition to ensuring that we only distribute products to licensed individuals. And they also wanted us to modify our suspicious order monitoring program to stop orders that we believed -- stop orders that could possibly be suspicious and then to any suspicious -- any order we deem suspicious should not be shipped.⁷⁴⁶

Although AmerisourceBergen began implementing these changes in 2007, the DEA, in fact, had given ABC notice two years earlier that its program was deficient, and these items needed improvement.

AmerisourceBergen met with the DEA in Washington, D.C. in August 2005. During that meeting, the DEA reminded AmerisourceBergen of several key compliance points including:

- Simply reporting suspicious orders does not relieve a distributor of the need to maintain effective controls against diversion;⁷⁴⁷
- The DEA cannot tell a distributor if an order is suspicious and so distributors must determine which orders are suspicious and make sales decisions;⁷⁴⁸
- Invalid prescriptions are not for legitimate medical needs and thus are diverted, regardless of where filled;⁷⁴⁹ and
- Any distributor “who is selling controlled substances that are being dispensed outside the course of professional practice must stop immediately.”⁷⁵⁰

Thus, as of 2005, the DEA placed ABC on notice that its current program was inadequate. However, like both McKesson and Cardinal Health, AmerisourceBergen simply did not implement the necessary changes. It was not until two years later when the settlement compelled them to do so that ABC began making substantive improvements in its anti-diversion program.

ABC’s controlled substances compliance program was outlined by several policy and procedure documents, which are listed at Appendix E, Figure 2.⁷⁵¹ [REDACTED]

⁷⁴⁶ See C. Zimmerman Deposition at 139:20-140:8.

⁷⁴⁷ See Meeting with AmerisourceBergen DEA Headquarters, *Internet Pharmacy Data*, 7 (Aug. 10, 2005), ABDCMDL00315887 [“DEA HQ Mtg.”].

⁷⁴⁸ See DEA HQ Mtg. at 8.

⁷⁴⁹ See DEA HQ Mtg. at 12.

⁷⁵⁰ See DEA HQ Mtg. at 13.

⁷⁵¹ See also AmerisourceBergen Diversion Control Program Policies & Procedures, ABDCMDL00003367 to ABDCMDL00003429; Email from D. May to C. Conneely, *et al.*, *Diversion Control Policies* (Jan. 15, 2015) (Caroline Conneely was with FTI), ABDCMDL00251385 to ABDCMDL00251406.

[REDACTED]

A simple review of the list in Appendix E, Figure 2 supports that finding. ABC's diversion control policies and procedures are contained within two different and separate series. One series starts with the prefix "DCP" and the other "CSRA." However, the documents are not formatted consistently across the series. For example, the CSRA series lists authors while the DCP series does not. Also, the DCP series consistently notes revision history while the CSRA series does so on an inconsistent basis. Finally, both series do not contain any linkage to a document's approval either by listing the names of the approvers or referencing a Document Change Notice ("DCN").

When taken together, both series contain documents covering the same topics, but containing different provisions, and there is no clear indication of how they link together. For example, there are two policies for the Order Monitoring Program (DCP-12.2.0 and CSRA 2.12). The purpose of DCP-12.2.0 states it is:

to establish the AmerisourceBergen Drug Corporation ("ABDC") Order Monitoring Program ("OMP") as a component of the broader Diversion Control Program, which is designed to prevent, detect and investigate the potential diversion of controlled substances and listed chemicals (hereafter referred to collectively as Controlled Substances) into other than legitimate medical, scientific and industrial channels. This policy establishes the requirement for ABDC's reviewing orders of Controlled Substances placed by ABDC customers in order to identify and investigate potentially suspicious orders and for reporting suspicious orders to the Drug Enforcement Administration ("DEA") and state authorities, as appropriate.⁷⁵³

The CSRA OMP policy's purpose states:

To ensure compliance with applicable state and federal regulations, AmerisourceBergen Corporation (ABC) has designed this program to review the ordering activity of its customers to identify possible excessive or suspicious orders of controlled substances and listed chemicals.⁷⁵⁴

Although the CSRA policy pre-dates the DCP policy (2005 versus 2007), the list of policy and procedure references in the DCP policy fail to list the CSRA policy even though the DCP policy seems to establish the mandate for the OMP.⁷⁵⁵ Overall, ABC has created unnecessary complexity through confusing sets of program documentation, which makes it hard to decipher exactly how ABC's program operated and for ABC to establish that their program indeed was effective.

⁷⁵² [REDACTED]

⁷⁵³ DCP-12.2.0 at § 1.1, ABDCMDL00003367 at ABDCMDL00003380.

⁷⁵⁴ See CSRA 2.12 at "Purpose."

⁷⁵⁵ DCP-12.2.0 at § 3, ABDCMDL00003367 at ABDCMDL00003380.



The diagram above outlines how AmerisourceBergen's order monitoring program ("OMP") operated beginning in 2007, showing the division of responsibilities between the Distribution Centers and CSRA.⁷⁵⁶

Beginning in 2007, ABC's OMP was based on the following parameters:

- [REDACTED]
- [REDACTED]
- [REDACTED]
- [REDACTED]
- [REDACTED]
- [REDACTED]

Despite AmerisourceBergen's efforts, including its perception that the DEA somehow endorsed it, the OMP had multiple inconsistencies and weaknesses.

A. Customer Size & Controlled Substances Ratio

[REDACTED] It is the interplay between these concepts and how CSRA applied them that ultimately resulted in the lack of a consistent, and hence effective, SOM and controlled substances program.

⁷⁵⁶ See Stephen Mays, *ABC Diversion Control Program effective June 25, 2007*, 18 (Jun. 25, 2007), ABDCMDL00000101.

⁷⁵⁷ See AmerisourceBergen, *Diversion Control Enhancements – Internal Update*, 6-7 (May 6, 2016), ABDCMDL00276831 ["DC Enhancements 2016"].

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

⁷⁶⁰ See Letter from J. Rannazzisi to All Registrants, 3 (Sep. 27, 2006).

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

The CSRA after-the-fact-reviews further undermined the reporting of suspicious orders and was not in compliance with the DEA's expectations communicated to ABC in 2005, because it reversed the paradigm that such orders needed to be investigated first, then shipped if the investigation found that they were not suspicious. In short, Mr. Zimmerman's proposal voided an established SOM control and abrogated CSRA's order monitoring oversight responsibilities.

C. OMP – Setting the Record Straight

In January 2012, ABC distributed an internal document entitled "Order Monitoring Program (OMP) Setting the Record Straight" which combined a high-level overview, frequently asked questions and talking points surrounding changes to OMP and deployment of SAP to manage ABC customer accounts.⁷⁷¹ The document, which was updated once more in October 2012, did anything but set the record straight.⁷⁷² Overall the October revisions simply "watered down" the controls around suspicious order monitoring program.

Both the January and October versions reference the fact that thresholds were now being applied by DEA registration number and not account number.⁷⁷³ Prior to the 2012 SAP upgrade, multiple ABC accounts could have the same DEA registration number.⁷⁷⁴ [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

⁷⁷⁶

The October version also removed the following language:

⁷⁷⁰ *Id.* (emphasis added).

⁷⁷¹ See Memorandum Order Monitoring Program (OMP) Setting the Record Straight, 1 (Jan. 30, 2012), ABDCMDL00002405. The document was sent under the auspices of Edward Hazewski. See Email from E. Hazewski to T. Cooper, Document (Jan. 16, 2010), ABDCMDL00280993 ["Setting the Record Straight Jan."].

⁷⁷² See Memorandum Order Monitoring Program (OMP) Setting the Record Straight (Oct. 3, 2012), ABDCMDL00280994 ["Setting the Record Straight Oct."].

⁷⁷³ See Setting the Record Straight Oct. at 5.

⁷⁷⁴ See Setting the Record Straight Jan. and Oct. at 5, FAQ #2.

⁷⁷⁵ See Setting the Record Straight Jan. and Oct. at 5, FAQ #2.

⁷⁷⁶ See E. Hazewski Deposition at 268:5-12.

[REDACTED]

The new version eliminated the use threshold review forms and the documented rationale for why the increase was needed.⁷⁷⁸ It also appears that blanket threshold increases across all drug families were now permitted.

The October version also eliminated FAQ #5 in the January version which stated:

OMP is based on order quantity, and it is meant to identify suspicious ordering patterns. The **DEA does not focus on what we ship to our customers**; they require us to evaluate what was ordered because the attempt to place an order is what can identify suspicious orders.⁷⁷⁹

Beyond being a simple misstatement of the law and regulations, the answer implies that if the order is not flagged as suspicious, ABC could adjust the shipment after the order cleared to accommodate customers. What is even more troubling is that this advice is being provided by the individual (Edward Hazewski) who headed up the Diversion Control team from 2008 to 2014.⁷⁸⁰

The October version of the communication professed that with the implementation of SAP, Distribution Center ("DC") discretion to release orders held due to OMP was being curtailed so that only "under limited circumstances" could the DC release held orders.⁷⁸¹ In reality, the October amendments allowed the DC to carry on releasing orders above the threshold.

According to the January [REDACTED]

[REDACTED]⁷⁸² After SAP, all DC discretion to release orders above the threshold was revoked.⁷⁸³ According to the January version, this was because:

[h]istorically, each DC had the ability to review held orders and apply their best judgment in releasing individual orders. Most sales associates have had accounts exceed their thresholds at some point in time; however, the DC had the ability to "make the call" after conducting their review which led to customers receiving their orders. As we deploy SAP to our DCs, the OMP management process becomes more systemic and less arbitrary. This is by design.⁷⁸⁴

⁷⁷⁷ See Setting the Record Straight Oct. at 5, FAQ #2 (emphasis in the original).

⁷⁷⁸ See Setting the Record Straight Oct. at 5, FAQ #3 (The October version eliminated the following sentence, "[t]he Account Manager can complete and submit a Threshold Review Form to ensure that this information [explaining the situation] is provided to Corporate Security and Regulatory Affairs.>").

⁷⁷⁹ See Setting the Record Straight Jan. at 5, FAQ #5.

⁷⁸⁰ See E. Hazewski Deposition at 21:5-12.

⁷⁸¹ See Setting the Record Straight Oct. at 7, FAQ #10.

⁷⁸² See Setting the Record Straight Jan. at 6-7, FAQ #10.

⁷⁸³ See Setting the Record Straight Jan. at 7, FAQ #10.

⁷⁸⁴ See Setting the Record Straight Jan. at 7, FAQ #10.

The October version removed the historical language above, as well as the requirement for a Threshold Review form to be filed with CSRA and allowed DC's to continue filling orders under the unspecific "limited circumstances" criterion.⁷⁸⁵ The net result of the October change was to dial back CSRA's visibility of the DC's threshold overrides and provided the DC's ability to approve overrides with less accountability. Therefore, while this change was good for sales, it negatively impacted the SOM program.

The October version also removed any language prohibiting disclosure to accounts that ABC was reporting rejected orders to the DEA as suspicious. In particular, the October version deleted the following sentence: "Notifying a customer that they have been reported to the DEA or State would defeat the purpose of the monitoring program."⁷⁸⁶ I can only conclude that ABC made the change to allow its sales force to provide customers with valuable coaching on how to avoid their orders being labeled as "suspicious," thereby undermining the SOM program even further. This is one of clearest instances that ABC cared more about preserving the customer relationship than exercising its statutory and regulatory obligations.

D. Low-Volume Accounts

In approximately July 2013, ABC began to work on a campaign to target its low-volume accounts. These were accounts with a low dollar volume (e.g., [REDACTED]), in other words, small retail pharmacies, but with a high ([REDACTED]) ratio of controlled substances.⁷⁸⁷ Internal sales talking points were developed using the threat that "your percentage of C2 orders is high and may be deemed suspicious by either our OMP system or regulatory authorities. This puts your account with ABDC at significant risk of closure or exposure to regulatory and enforcement agencies actions."⁷⁸⁸

The goal was not to lower the ratio of controlled substances purchases, but rather to have the pharmacy "make ABDC your primary wholesaler and shift all purchases to us [ABDC]" because "[i]ndependent pharmacies come to AmerisourceBergen because they know that by taking full advantage of our product offerings, resources, and expertise, they will be able to attract more patients, retain existing patients and improve their operating efficiencies."⁷⁸⁹ In other words, AmerisourceBergen would continue supplying these pharmacies with controlled substances at their usual levels provided they could get the controlled substances ratio down by purchasing a more non-controlled product from ABC. Neither Messrs. Zimmerman nor Mays in their depositions could provide an alternative rationale for the document.

Setting aside the clearly unethical stance of using potential compliance violations to intimidate customers, AmerisourceBergen was not engaging in this program because it was concerned about the diversion of controlled substances, as it publicly professed, but rather ABC was engaging in a good, old fashioned switch program using a negative incentive in place of the more commonly used positive ones. ABC was attempting to increase its market share among the small retail pharmacies while maintaining and increasing its profits.

⁷⁸⁵ See Setting the Record Straight Oct. at 7, FAQ #10.

⁷⁸⁶ See Setting the Record Straight Oct. at 7, FAQ #11.

⁷⁸⁷ See C. Zimmerman memorandum to E. Hazewski, *et al.*, *RVP Talking Points*, 1 (Jan. 19, 2009), ABDCMDL00000169.

⁷⁸⁸ See Sales Talking Points, Low-Volume Accounts (Jul. 2013), ABDCMDL00278212.

⁷⁸⁹ See Sales Talking Points, Low-Volume Accounts (Jul. 2013), ABDCMDL00278212.

Once more, in August 2017, ABC re-visited the problem with customers utilizing AmerisourceBergen as a secondary distributor for purchasing controlled substances. ABC looked at preventing customers from purchasing four high-risk drug families (including hydrocodone solid, hydromorphone solid, oxycodone 30 mg solid and oxycodone solid) from ABC as a secondary distributor.⁷⁹⁰ ABC analyzed the impact of this potential alteration to its program from the perspective of dosage units and dollar volume. The change in its business would have resulted in a decrease of █████ dosage units of just the four drug families considered – roughly █████ of the company’s business in those four drug families.⁷⁹¹ The financial impact to ABC’s total controlled substance sales by removing four drug families included a loss of almost █████ or roughly █████ of ABC’s total controlled substance sales.⁷⁹² Predictably, ABC chose not limit the ability for its customers to purchase these four drug families from ABC as a secondary supplier.⁷⁹³

However, ABC, with the help of its own compliance staff, undermined its own SOM program by providing a “work around” to the product mix control. For example, a top purchaser review of █████ was conducted in January 2010, three years after █████ threshold for benzodiazepine anxiety solids was set at █████ dosage units based on a threshold override in 2007.⁷⁹⁴ While the review noted that █████ was a medium volume, high controlled substance ratio account (█████ average monthly dollar volume and a █████ controlled substances ratio), the review simply noted that the █████ purchases had not reached the threshold in the past two years after having been raised and simply requested that the CSRA 590 be completed.⁷⁹⁵

11.5.3 AmerisourceBergen’s post-settlement customer due diligence program was ineffective as a result of poor design and inconsistent application.

Enhanced customer due diligence was one of the major improvements that the 2007 DEA-ABC settlement was intended to achieve.⁷⁹⁶ Yet, in reality, while ABC did establish a “policy” or “process” governing customer account due diligence in May 2007,⁷⁹⁷ CSRA did not require centralized customer files until six years later in 2013⁷⁹⁸ and did not create implementing procedures until a full ten years later in 2017.⁷⁹⁹

⁷⁹⁰ See Email from M. Guerreiro to D. May RE: Secondary Account Review, (Aug. 18, 2017), ABDCMDL00354768; Presentation by ABC, Controlled Substances Limitation Analysis, 2 (undated), ABDCMDL00354771); ABC Chart, OMP of Interest, ABDCMDL00354777.

⁷⁹¹ See *id.* at 3; see also Marcellino Guerreiro Deposition, 278:10-284:8 (April 3, 2019).

⁷⁹² *Id.*

⁷⁹³ *Id.*; see also M Guerreiro Deposition at 284:22.

⁷⁹⁴ See Memorandum from K. Kreutzer, *Top Purchaser Review – BSD Inc.* (Jan. 11, 2010), ABDCMDL00158760 [“BSD Top Purchaser Review”].

⁷⁹⁵ See BSD Top Purchaser Review.

⁷⁹⁶ See C. Zimmerman Deposition at 139:20-140:8.

⁷⁹⁷ See Amerisource Bergen, Policy CSRA 3.4 Customer Account Due Diligence, (Feb. 13, 2013) (The original effective date was May 8, 2007), ABDCMDL00251385 at ABDCMDL00251400.

⁷⁹⁸ See AmerisourceBergen, Policy CSRA 3.5 Customer Due Diligence Documentation (May 10, 2013) (“This is a new policy.”), ABDCMDL00251385 at ABDCMDL00251402.

⁷⁹⁹ See AmerisourceBergen, Procedure DCP SOP-12.1.10 (Jan. 1, 2017).

A. The CSRA Form 590

The 2007 Account Due Diligence Policy was intended “[t]o establish a process of due diligence and on-going screening of New Customer Accounts to ensure AmerisourceBergen (ABC) only services Customer Accounts that comply with Federal and State Regulations as well as ABC Policy.”⁸⁰⁰ The same document, however, states “that the following policy applies ...” and that by not distinguishing between a policy and an SOP, authors Messrs. Mays and Hazewski merely confused the situation.⁸⁰¹

The “process” was to complete the CSRA Form 590 (the retail pharmacy questionnaire), next to have an ABC Business Development Manager (“BDM”) conduct a site visit, and then have CSRA “verify” the information and either approve or deny the request.⁸⁰² The Account Due Diligence policy mandated that the CSRA 590 was to be completed by the BDM during the site visit.⁸⁰³ Although originally established for new customers, sometime after 2007, ABC began using the CSRA 590 with existing customers, especially those with expanding or changing business models.⁸⁰⁴ However, retail chain pharmacies, defined as pharmacies having more than ten stores, were exempt from having a Form 590 on file.⁸⁰⁵

The importance that ABC placed on the CSRA Form 590 as a critical anti-diversion control was substantial. For example, the CSRA Form 590 itself stressed that the “questionnaire is an official business record subject to review by State and Federal regulatory agencies and stipulated that “[f]orms that are not **complete**, illegible, or not signed will be returned.”⁸⁰⁶

As ABC told the House Energy and Commerce Committee:

The information contained on the questionnaire is the basis for ABDC’s due diligence investigation and provides a baseline to measure the pharmacy’s ordering habits and to determine any deviation from expected purchasing practices. The questionnaire provides information to ABDC regarding anticipated ordering practices, including, among other things, the amount of controlled substances ordered, the anticipated ratio of controlled vs. non-controlled substances purchased, key prescribing doctors in the area utilizing the pharmacy, the purchasing practices of the pharmacy’s customers (i.e. cash, credit, insurance, etc.), and whether another supplier is known to have suspended or ceased controlled substance sales to the customer. The questionnaire also includes inquiries on topics such as high-risk drugs and high-prescribing physicians.”⁸⁰⁷

⁸⁰⁰ See Account Due Diligence, Policy CSRA 3.4 at “Purpose”.

⁸⁰¹ See Account Due Diligence, Policy CSRA 3.4 at “Purpose”.

⁸⁰² See generally Account Due Diligence, Policy CSRA 3.4.

⁸⁰³ See Account Due Diligence, Policy CSRA 3.4 at “A. Retail Pharmacy Questionnaire”.

⁸⁰⁴ See C. Zimmerman Deposition at 325:2-12.

⁸⁰⁵ See C. Zimmerman Deposition at 213:16-214:22.

⁸⁰⁶ See ABC Form CSRA-590, Information (Apr. 2015) (emphasis added), ABDCMDL00355073; see also N. Elkins Deposition at 228:6-10 (confirming the CSRA Form 590 information was “vital”).

⁸⁰⁷ See W.Va. Red Flags Report at 113-114.

B. “Verifying” CSRA 590 Data

Once the site visit was completed and the CSRA 590 filled out, it became CSRA’s task to verify the information provided.⁸⁰⁸ CSRA verification was limited to reviewing the questionnaire responses, performing internet searches on the information provided, and verifying the customer was not listed on ABC’s Do Not Ship List.⁸⁰⁹ CSRA verification was limited to using a checklist with a series of “YES/NO” responses with extremely limited space to add pertinent details, and which did not instruct the completer that additional information could be appended.⁸¹⁰ Consequently, the verification performed by CSRA was of little or no utility in determining whether to approve the customer.⁸¹¹

Furthermore, while ABC’s description of the CSRA Form 590 to the House Energy and Commerce Committee technically was accurate, it neglected to mention that neither the Account Due Diligence Policy or the Form 590 required supporting source documentation be provided concerning pharmacy ordering and dispensing practices, as well as key prescriber habits. Therefore, CSRA could not independently verify these critical indicators of potential diversionary customer behavior rendering the CSRA verification step simply a “pro forma” exercise.

As ABC told the House Energy & Commerce Committee, “ABDC does, at times, request dispensing data from both current and prospective customers. There is no specific frequency at which dispensing data is requested from customers.”⁸¹² ABC further elaborated that it “collects patient de-identified dispensing reports on an as-needed basis to allow it to investigate and mitigate concerns about possible suspicious behavior by its customers[,]” and that “[c]ustomers may also be asked to provide full dispensing reports as part of new customer due diligence, again to mitigate red flags discovered during onboarding or to properly size the pharmacy as part of the company’s Ordering Monitoring Program.”⁸¹³ When pressed further about why it did not routinely request this data, ABC told the Committee “[c]ollecting dispensing data on a routine basis from all pharmacies is not a requirement that is imposed upon the distributor by the governing federal laws and implementing regulations.”⁸¹⁴

⁸⁰⁸ See Account Due Diligence, Policy CSRA 3.4 at “D. Account Verification Checklist”.

⁸⁰⁹ See Account Due Diligence, Policy CSRA 3.4 at “D. Account Verification Checklist”. CSRA used the CSRA Form 590c to perform the verification. See Retail Pharmacy Questionnaire (undated), ABDCMDL00301401; see also Customer Verification Checklist (Feb. 9, 2010), ABDCMDL00288989; see also Email from E. Cherveny to E. Coldren, *et al.*, FYI: OY Block – Pharmlink, Inc. (10056420/BP6335156), (Sep. 22, 2016), ABDCMDL00246107-109; Email from E. Coldren to M. Guerraio, FW: DNS List, (Feb. 21, 2018), ABDCMDL00300163-166. Recent examples of ABC’s failure to enforce the “Do Not Ship” List.

⁸¹⁰ See Retail Pharmacy Questionnaire (undated), ABDCMDL00301401; see also Customer Verification Checklist (Feb. 9, 2010), ABDCMDL00288989.

⁸¹¹ See Email from D. May to S. Hartman, *et al.*, RE: Mingo Pharmacy, (Oct. 11, 2017), ABDCMDL00142299. ABC failed to recognize that a customer gave incorrect information on its 590 form until ABC was notified by DEA during an investigation of the customer. However, even then ABC did not terminate the customer.

⁸¹² See W.Va. Red Flags Report at 114 (citing *Combating the Opioid Epidemic: Examining Concerns About Distribution and Diversion: Hearing Before the Subcomm. on Oversight and Investigations of the H. Comm. on Energy and Commerce*, 115th Cong. (2018), Responses to Questions for the Record submitted by Steven H. Collis, CEO, President and Chairman of the Board, AmerisourceBergen Corp.).

⁸¹³ See W.Va. Red Flags Report at 114.

⁸¹⁴ See W.Va. Red Flags Report at 114.

Although ABC's counsel is correct that neither the CSA nor its implementing regulations explicitly require ABC to collect this data, ABC's failure to do so effectively undermined the entire due diligence process rendering it moot. Without independent dispensing reports, CSRA was limited to "verifying that [listed] physicians are in good standing to prescribe."⁸¹⁵ Even that control could be circumvented easily by providing CSRA with a list of physicians in good standing, regardless of whether they were the pharmacy's top prescribers of controlled substances.

However, the flawed process design did not end there. The Account Due Diligence Policy also did not clearly define verification roles and responsibilities within CSRA, nor did it specify the criteria for determining whether to accept or reject an account.

C. CSRA 590 Validation Project

In 2016, the Diversion Control Team began a review project of the CSRA 590s in 2016. The purpose of this project was "to validate that all current ABDC customers authorized to purchase controlled substances have the required due diligence documentation in the file."⁸¹⁶ In its first phase, the "project was to conduct a full review of every ABDC customer authorized to purchase controlled substances and identify any with deficiencies."⁸¹⁷

[REDACTED]

[REDACTED]

[REDACTED]

D. Process Not Followed

The CSRA Form 590 Due Diligence program is another example of the differing public face and private face of ABC's compliance efforts. Publicly, AmerisourceBergen touted the Form 590s as its improved due diligence effort. [REDACTED]

This was further compounded by the fact that ABC simply did not follow its declared due diligence process. The CSRA 590 Validation Project shows that ABC was not following the due diligence process established in 2007 in response to its settlement with the DEA. With 3,285 customers having no CSRA 590, ABC clearly did not have adequate controls in place to ensure that the CSRA 590s were complete before approving a new account or as a reference on existing customers. Nor were ABC's controls adequate for the CSRA 590 to be used as a reliable source of information when pharmacy circumstances changed.

ABC also lacked an adequate corrective action process to remedy the large shortfall, as demonstrated by the fact that almost 12 months after identifying the "sheer volume" of customers with incomplete or missing CSRA 590s,⁸²⁰ ABC had managed to bring about [REDACTED].⁸²¹

⁸¹⁵ See C. Zimmerman Deposition at 343:3-6.

⁸¹⁶ See Email from E. Cherveney to S. Hendrickson, *et al.*, CSRA Validation Project (Aug. 5, 2016), ABDCMDL00159415.

⁸¹⁷ *Id.*

⁸¹⁸ *Id.*

⁸¹⁹ See CSRA Form 590 Validation Project Spreadsheet (Jul. 28, 2016), ABDCMDL00159417.

⁸²⁰ See Email from E. Cherveney to S. Hendrickson, *et al.*, CSRA Validation Project (Aug. 5, 2016), ABDCMDL00159415.

Therefore, the compliance efforts that ABC espoused publicly as important were not made a priority within the company.

11.5.4 Like its order monitoring and customer due diligence processes, ABC's investigations process suffered from poor design and lack of consistency rendering it ineffective.

Although the due diligence and investigations process was rolled out in June 2007,⁸²² it was more than a year later in October 2008, that CSRA implemented a policy governing targeted site visits (i.e., investigations).⁸²³ CSRA created the policy to address "specific concerns found during the course of data review or information received from outside sources."⁸²⁴ According to the Targeted Visits policy, "CSRA will identify customer locations to be visited based on factors, including but not limited to; Diversion Control Program data analysis; information from ABC personnel, regulatory agencies, or any other sources; and/or at the request of [the] ABC General Counsel."⁸²⁵ However, it was not routine practice to perform targeted visits on existing customers.⁸²⁶

Like other CSRA policies, the Targeted Visits policy failed to outline, except in the most general terms, the factors CSRA used to trigger an onsite visit. Some of the unwritten specific criteria that could trigger a targeted visit involved "the activity of the customer, whether they wanted changes, whether they were, you know, being changed in the program, whether they were increasing sales or changing areas of service."⁸²⁷

According to the Targeted Visits policy, the Director or Program Manager of the Diversion control program would "select the CSRA Representative or contractor to conduct the visit based on availability and their proximity to the location in question."⁸²⁸ In addition to the dedicated investigators and the members of the Diversion Control Team, CSRA also pulled in Distribution Center compliance staff to support investigations.⁸²⁹ The Director or Program Manager, however, was not required to assure the CSRA Representative or contractor was qualified.

Targeted visits typically were announced visits of limited duration. Although, the policy states that "[s]urveillance visits with no contact of pharmacy personnel may be conducted un-announced," the clear

⁸²¹ See Email from D. May to J. Sharkey, *et al.*, FW: CSRA 590 Validation Project (Jul. 7, 2017) [REDACTED], ABDCMDL00159415.

⁸²² See C. Zimmerman Deposition at 211:9-12.

⁸²³ See Amerisource Bergen, Policy CSRA 2.25 Retail Pharmacy Targeted Visits, (Jun. 17, 2013) (The original policy became effective on Oct. 1, 2008), ABDCMDL00251385 at ABDCMDL00251392.

⁸²⁴ See Targeted Visits, Policy CSRA 2.25 at "Purpose".

⁸²⁵ See Targeted Visits, Policy CSRA 2.25 at "Policy".

⁸²⁶ See C. Zimmerman Deposition at 318:1-4 and 321:4-322:5.

⁸²⁷ See C. Zimmerman Deposition at 209:17-22.

⁸²⁸ See Targeted Visits, Policy CSRA 2.25 at "Overview of Procedure".

⁸²⁹ See C. Zimmerman Deposition at 449:17-451:8.

expectation was that targeted visits required scheduling.⁸³⁰ Therefore, the typical visit required CSRA to contact the Account Manager to coordinate the visit.⁸³¹ The Account Manager then contacted “the owner/pharmacist in charge (PIC), inform him/her of the visit, and ensure that the owner/PIC or their designee will be present on the arranged date in order to give their **undivided attention to the CSRA representative for a minimum of one hour.**”⁸³² The implication was that any fieldwork at the site would be completed in an hour.

Although the policy mandated that the investigator must prepare a summary report “to include all observations or concerns noted during the visit as well as recommendations provided to the DC for correction of deficiencies” and “follow a memo format” [REDACTED].⁸³³

For example, in October 2013, [REDACTED]⁸³⁴ In her email report to [REDACTED]

[REDACTED]
[REDACTED]
[REDACTED]
[REDACTED]⁸³⁶

CSRA investigations, like the verifications of Form 590, were at best cursory. For example, the House Energy and Commerce Committee noted that in December 2015 ABC failed to investigate Westside Pharmacy before reinstating the pharmacy as a customer.⁸³⁷ According to the Committee, Westside told ABC that its ability to purchase controlled substances had been either terminated or restricted by a distributor in the past, and the CSRA 590 was provided to ABC on the same day it was terminated by Miami-Luken in response to a DEA Show Cause Order.⁸³⁸ Furthermore, two of the top five listed prescribers of hydrocodone and oxycodone were the subject of public enforcement actions or had a DEA license that could not be verified.⁸³⁹ ABC’s response was that “[n]ews searches for prescribing physicians are not a standard part of ABC’s new customer review.”⁸⁴⁰

⁸³⁰ See Targeted Visits, Policy CSRA 2.25 at “1-Notifications”.

⁸³¹ See *id.* at “1-Notifications”.

⁸³² See Targeted Visits, Policy CSRA 2.25 at “2 Pre-Visit Preparation” (emphasis added).

⁸³³ See Targeted Visits, Policy CSRA 2.25 at “6-Reports”.

⁸³⁴ See Email from T. Polster to R. Swords, ABC Visit (Oct. 31, 2013), WAGMDL00237263.

⁸³⁵ See *id.*

⁸³⁶ See *id.*

⁸³⁷ See W. Va. Red Flags Report at 163.

⁸³⁸ *Id.*

⁸³⁹ See *id.* at 163-166.

⁸⁴⁰ *Id.* at 166.

manufacturing partners, and customer/buying groups,”⁸⁴⁷ the company’s approach to controlled substances training can be best described as “scattershot.”

While ABC did attempt to provide training to the primary groups involved with SOM and controlled substances such as the sales teams, compliance managers, and even CSRA staff members, its training efforts were both fragmented and inconsistent. On the one hand, ABC, in the course of discovery in this case, produced several slide decks supposedly used as part of various training sessions.⁸⁴⁸ On the other hand, no corresponding “sign-in” sheets or other records were located to support when specific sessions occurred and who attended, despite Mr. Zimmerman’s 2014 inquiry to Cathy Marcum who responded that training was done “[a]nnually and it is documented on paper training logs.”⁸⁴⁹

The result is that although Ed Hazewski testified that “everyone in the organization had some exposure to discussions and training that related to potential diversion issues,”⁸⁵⁰ Stephen Mays could not recall if ABC required or maintained any records that would support Mr. Hazewski’s contention.⁸⁵¹ Mr. Hazewski’s contention was also contradicted by ABC’s own employees.

For example, Julie Fuller, an ABC sales representative, stated that as an account manager, ABC only provided her general sales training, and did not provide any training or information on “(a) how to identify questionable pharmacy behavior like suspicious dispensing, sales, or prescription filling practices, (b) how to report concerns regarding those behaviors, or (c) how to ensure that account managers only signed up and maintained accounts with legitimate pharmacies.”⁸⁵² This is consistent with what Mr. Zimmerman found in 2012 when he wrote to Mr. Hazewski that “[s]o far we have not met anyone from sales, VPs, RVPs, Director, Specialists, etc., that is aware of the OMP training, are you sure this has been rolled out and communicated?”⁸⁵³ Later he wrote to Mr. Hazewski, “[w]e should work with Sales to ensure we know [who] is going to monitor and track ... to make sure all sales folks go through the training.”⁸⁵⁴

Later in 2014, Greg Madsen noted that neither the OMP policy nor the training were current, writing to Stephen Mays that:

⁸⁴⁷ See ABC, *AmerisourceBergen Diversion Control Program Overview*, 3 (Jan. 2016) (“Education and Training” section), ABDCMDL00303979 [“Diversion Control Program Overview”]

⁸⁴⁸ See, e.g., ABC Presentation, *Diversion Control Training Program for Sales Associates*, (Jul. 30 to Aug. 2, 2014) (The meeting was held at the MGM Grand Hotel in Las Vegas, Nevada), ABDCMDL00158342; ABC Presentation, *Diversion Control Program & OMP – General Awareness Training*, (Apr. 2016); ABC Presentation, *OMP Strategy for Retail Accounts – Low Dollar Controlled Substances High Risk*, (undated), ABDCMDL00282234; ABC Presentation, *Prescription Drug Diversion – Recognizing the Red Flags*, (undated), ABDCMDL00269475; ABC Presentation, *Diversion Control Program – DC OMP Training*, (May 2009), ABDCMDL00141716.

⁸⁴⁹ See Email from C. Marcum to C. Zimmerman, *et al.*, Re: DCP/OMP Training Questions, (Feb. 27, 2014), ABDCMDL00277674.

⁸⁵⁰ See E. Hazewski Deposition at 44:16-19.

⁸⁵¹ See S. Mays Deposition at 239:19-21.

⁸⁵² See Declaration of Julie Fuller ¶9 (Jan. 26, 2019).

⁸⁵³ See Email from C. Zimmerman to E. Hazewski, *et al.*, Fwd: CSRA Training – Questions, (Dec. 20, 2012, 11:17 AM), ABDCMDL00269150.

⁸⁵⁴ See Email from C. Zimmerman to E. Hazewski, *et al.*, Re: CSRA Training – Questions, (Dec. 20, 2012, 11:17 AM), ABDCMDL00269150.

[REDACTED]

Mr. Mays simply responded, [REDACTED].⁸⁵⁶

However, “soon” for Amerisource Bergen meant two years later. A string of emails authored in January 2016 show that despite expressing a recognition that OMP training needs to be conducted annually, ABC persisted in using an OMP training module described by various different employees as “ancient history,” and “sorely outdated.”⁸⁵⁷ Furthermore, in the absence of an established CSRA program, the distribution centers created their own OMP training “which was all over the map” in an effort to fill the gap.⁸⁵⁸

Contributing to this fragmented and inconsistent approach was the fact that CSRA had no formal, documented training process for controlled substances compliance that was located. At a minimum, this training process should have detailed the mandatory training curriculum, the maintenance of training records, and how completion is monitored. However, it does not appear to exist in either the Diversion Control Program or the CSRA policies and procedures pertaining to controlled substances.⁸⁵⁹ Therefore, ABC’s claims of mandatory training in the Diversion Control Program Overview could not be verified by me.⁸⁶⁰

CSRA also did not have specific staff members directly responsible for implementing, overseeing and monitoring training. Instead CSRA apparently opted for making the entire team responsible for training.⁸⁶¹ As a result, FTI concluded that:

[REDACTED]⁸⁶²

⁸⁵⁵ See Email from G. Madsen to S. Mays, RE: DCP/OMP Training Questions, (Feb. 27, 2014, 2:10 PM), ABDCMDL00291500.

⁸⁵⁶ See Email from S. Mays to G. Madsen, RE: DCP/OMP Training Questions, (Feb. 27, 2014, 2:10 PM), ABDCMDL00291500.

⁸⁵⁷ See Email from E. Cherveney to N. Seckinger, RE: OMP-General Training for all Associates (Jan. 15, 2016), ABDCMDL00151814.

⁸⁵⁸ See Email from E. Cherveney to N. Seckinger, RE: OMP-General Training for all Associates (Jan. 15, 2016), ABDCMDL00151814.

⁸⁵⁹ See Appendix E, Figure 2.

⁸⁶⁰ See Diversion Control Program Overview at 3 (“Education and Training” section).

⁸⁶¹ See S. Mays Deposition at 239:6-7.

⁸⁶² See FTI Narrative at 8.

Consequently, FTI concluded that [REDACTED]

[REDACTED]⁸⁶³

[REDACTED]⁸⁶⁵

At some point after FTI's report, it appears that CSRA created an informal CSRA Training Committee. The first references to the Committee, however, do not appear until June 2016 indicating that it was a late addition to ABC's controlled substances compliance efforts.⁸⁶⁶ With the appearance of the CSRA Training Committee in June 2016 also comes the first reference to a controlled substances course list.⁸⁶⁷ However, the course list from June 2016 shows that many of the CSRA training programs, beyond just controlled substances, either were not being addressed or were incomplete.⁸⁶⁸

For any compliance function, even a foundational one, the failure to address these basic training issues represent at best simple sloppiness. In the context of a controlled substances compliance program, they are enough to conclude that by failing to adequately train its employees, ABC was not maintaining an effective anti-diversion program.

11.5.7 ABC's failure to implement a formal corrective action program contributed to the company's continued ineffective controlled substances compliance program.

Although ABC made periodic changes to its controlled substances compliance program during the review period, there is little evidence to suggest that these changes originated through an internal corrective action program. Rather, the impetus for most of the significant program changes come from external sources - the DEA and FTI.

It also is clear that ABC did not have a robust internal audit function that addressed controlled substances. In 2009, Mr. Hazewski sent a formal memorandum to Mr. Zimmerman discussing the findings from an audit of the

⁸⁶³ *Id.*

⁸⁶⁴ See D. May Deposition at 128:17-20.

⁸⁶⁵ See Email G. Madsen to G. Guevara and P. Neipp, FW: Lead Clerk Training, (Mar. 1, 2016) (emphasis added), ABDCMDL00311966.

⁸⁶⁶ See Email from F. Duncan to G. Crowley, *et al.*, CSRA Training Committee Weekly Meeting – Notes, Updated Course List & Actions attached, (Jun. 22, 2016), ABDCMDL00303935.

⁸⁶⁷ See Email from F. Duncan to G. Crowley, *et al.*, CSRA Training Committee Weekly Meeting – Notes, Updated Course List & Actions attached, (Jun. 22, 2016), ABDCMDL00303935.

⁸⁶⁸ See CSRA Training Program Overview Spreadsheet, (undated) (Entries date to early June 2016), ABDCMDL00303937; *see also*, K. Kreutzer email to J. Sutherland, *et al.*, FW: CSRA Training Committee – OMP Program Follow-up, (Jul. 20, 2016), ABDCMDL00303977.

OMP program conducted by Michael Mapes, an ex-DEA employee.⁸⁶⁹ To conduct his “audits,” Mr. Mapes utilized a checklist that cataloged the requirements from CSRA’s diversion control policies and procedures and noted “compliant” or “not compliant.”⁸⁷⁰ However, the checklist’s format does not allow the auditor to add details, explanations or even the samples that were pulled to support the compliant/not compliant designation.

Although billed as an “audit,” the Mapes Checklist is merely a quality control device. It certainly is not an “audit” that is designed to detect and highlight issues to be corrected. For example, in the case of CSRA Form 590, the checklist only inquires whether the Form 590 has been completed for all new retail pharmacy applicants and whether the 590c was being used by CSRA to verify responses.⁸⁷¹ Despite these questions being asked on an annual review of CSRA files, it appears that this “audit” program failed to uncover the fact that CSRA frequently was failing to require completed Form 590s until more than 3,200 customers were impacted. Nor did this “audit program” take issue with the fact that almost 12 months later, ABC was struggling to complete remediation on 10% of the outliers.

11.6 Accountability - Consistent Enforcement

11.6.1 AmerisourceBergen does not enforce the standards of the program, and thus there is no real accountability for the program’s lack of effectiveness.

AmerisourceBergen did not take steps to “remove” those individuals from positions of substantial authority for controlled substances compliance even though their actions and failures to act helped render the controlled substances compliance program ineffective to detect and prevent diversion. As a result, these individuals have yet to be held accountable for their roles in the controlled substances compliance program’s failure:

1. **Christopher Zimmerman** – Although recently removed from his position as ABC’s Chief Compliance Officer, Mr. Zimmerman remains in control of CSRA and thus AmerisourceBergen’s anti-diversion efforts.
2. **Stephen Mays** – The scope of Mr. Mays role within ABC’s diversion control program has varied over the years, but as head of ABC’s Pharmaceutical Distribution and Global Sourcing Group, he retains substantial sway over the implementation of the program.
3. **David May** – As the current head of Security and Diversion Control, Mr. May continues to have direct day-to-day oversight of ABC’s program.

The same lack of accountability can be seen with ABC’s sales representatives, As Nathan Elkins testified, no sales representative who worked for him was ever fired for not reporting a suspicious order.⁸⁷²

⁸⁶⁹ See Memorandum from E. Hawzewski to C. Zimmerman, *OMP Audit*, (Dec. 17, 2009).

⁸⁷⁰ See AmerisourceBergen, *Diversion Control Program Audit Checklist*, (Dec. 14, 2007) (The checklist notes that it is for internal corporate audits).

⁸⁷¹ See *id.* at 3, Questions 1 and 3.

⁸⁷² See N. Elkins Deposition at 158:8-15.

By failing to hold individuals accountable for the controlled substances program's established lack of effectiveness or for not complying with ABC's policies and procedures, AmerisourceBergen's corporate compliance program was mere words on paper that professed to hold culpable individuals accountable.

12 CVS Health Inc.

12.1 Background

CVS Health, Inc. ("CVS") began in 1963 as a single store selling health and beauty products in Lowell, Massachusetts by the brothers Goldstein (Stanley and Sidney) and their partner Ralph Hoagland.⁸⁷³ At the end of 2017, CVS, which stands for Consumer Value Stores, had more than 9,800 pharmacies in the U.S., Puerto Rico and Brazil employing more than 246,000 people in the U.S.,⁸⁷⁴ and net annual revenues of \$184.8 billion.⁸⁷⁵ Much of CVS' growth was accomplished by a series of strategic mergers including Caremark Rx, Inc. (PBM services) in 2007, Omnicare (pharmacies in long-term care facilities) in 2015, and Aetna (health insurance) in 2018.⁸⁷⁶

From 2006 to 2014, CVS distributed opioids (hydrocodone combination products or HCPs) from two distribution centers to its retail stores organized under the CVS Pharmacy, Inc. umbrella.⁸⁷⁷ The primary controlled substances distribution center in this case was CVS Indiana L.L.C. located in Indianapolis, Indiana and an ancillary center, CVS Rx Services LLC., located in Chemung, New York. With the reclassification of hydrocodone combination products from Schedule III to Schedule II in 2014,⁸⁷⁸ CVS ceased the internal distribution of any opioid products to its retail locations at the end of September that year.⁸⁷⁹

12.2 Executive Summary

CVS likes to describe itself as "a health care innovation company helping people on their path to better health ... transforming the consumer health care experience and helping to foster healthier communities."⁸⁸⁰ The

⁸⁷³ See CVS HEALTH, *History*, <https://cvshealth.com/about/company-history> (last visited Feb. 17, 2019).

⁸⁷⁴ See CVS HEALTH, *CVS Health at a Glance*, <https://cvshealth.com/about/facts-and-company-information> (last visited Feb. 17, 2019).

⁸⁷⁵ See CVS HEALTH, *Financial Highlights*, <https://investors.cvshealth.com/investors/2017-in-review/default.aspx> (last visited Feb. 17, 2019).

⁸⁷⁶ See CVS HEALTH, *History*, <https://cvshealth.com/about/company-history> (last visited Feb. 17, 2019).

⁸⁷⁷ For Schedule II opioids, CVS utilized other distributors including primarily the G1 distributors.

⁸⁷⁸ See 79 Fed. Reg. 49661 (Aug. 22, 2014).

⁸⁷⁹ See Email from D. Coakley, RE: Hydro Transition Meeting Questions, (Sep. 23, 2014) (referencing September 26, 2014 as the last shipment date for HCPs), CVS-MDLT1-000002195.

⁸⁸⁰ See CVS HEALTH, *Investor Story*, <https://investors.cvshealth.com/investors/investor-story/default.aspx> (last visited Feb. 17, 2019).

company's compliance actions, however, suggest that CVS does not view compliance as an important part of its efforts to transform the health care experience.

Overall, from 2008 to 2018, CVS has settled six major cases with the Justice Department, the HHS OIG, and the DEA. Half of the settlements involved health care fraud issues, including federal false claims allegations.⁸⁸¹ The other half involved repeated violations of the Controlled Substances Act and its accompanying regulations.⁸⁸² This is an exceedingly poor compliance record for one of the largest retail pharmacy chains in the U.S.

Coupled with its poor compliance record, CVS simply considered distribution center compliance with the controlled substances requirements to be an afterthought, and as a result, the DC anti-diversion program did not impact CVS' business or cultural equations in a meaningful way.

This lack of compliance prioritization and commitment manifested itself in four key respects:

- **Dual Roles:** Even though CVS simultaneously occupied both the dispenser and distributor roles for controlled substances in the "closed loop" system, CVS made little effort to incorporate its own pharmacy dispensing data within its SOM program.⁸⁸³ When CVS did make an effort to do so, the data used often was incomplete or "stale." The fact that CVS did not effectively do so was a missed opportunity to improve its program significantly.
- **Corporate Entity Structure:** The CVS organization has evolved an extremely complicated and compartmentalized corporate entity structure that is akin to the entity structure typically seen in Chinese or Japanese pharmaceutical companies. CVS appears to use that structure to avoid taking responsibility for obligations it wishes to avoid. In the case of the distribution center anti-diversion program, this structure, which created isolation and fragmentation, impeded any improvement efforts. As a result, CVS essentially shunned the program and avoided meeting its obligations in a forthright manner.
- **Failure to Adequately Prioritize Controlled Substances Compliance:** CVS prioritized revenue generation and preventing any disruption to the retail pharmacies' ability to fill prescriptions over any

⁸⁸¹ See Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Caremark Corporation (Mar. 14, 2008) https://oig.hhs.gov/fraud/cia/agreements/cvs_cia_executed.pdf; Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Caremark Corporation (Mar. 25, 2014), https://oig.hhs.gov/fraud/cia/agreements/ CVS_Caremark_03252014.pdf; Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Health Corporation (Oct. 11, 2016), https://oig.hhs.gov/fraud/cia/agreements/ CVS_Health_Corporation_10112016.pdf.

⁸⁸² Press Release CVS/pharmacy, *Lapse in controls of PSE sales in certain CVS/pharmacy stores in 2007 and 2008 relates to electronic monitoring system flaw that has been corrected Settlement amount fully reserved and previously disclosed; should have no further effect on company's financial results*, (Oct. 4, 2010), <https://cvshealth.com/newsroom/press-releases/cvsparmacy-announces-agreements-us-drug-enforcement-administration-and-us-attorneys-offices>; Press Release, U.S. Dep.'t of Justice, Drug Enforcement Admin., *CVS To Pay \$11 Million To Settle Civil Penalty Claims Involving Violations Of Controlled Substances Act*, (Apr. 3, 2013), <https://www.dea.gov/press-releases/2013/04/03/cvs-pay-11-million-settle-civil-penalty-claims-involving-violations-0>; Press Release, U.S. Dep.'t of Justice, Drug Enforcement Admin., *CVS Pharmacy, Inc. to pay \$1.5 million to settle Civil Penalty Claims for violation so [sic.] the Controlled Substance Act*, (Jun. 28, 2018), <https://www.dea.gov/press-releases/2018/06/28/cvs-pharmacy-inc-pay-15-million-settle-civil-penalty-claims-violation-so>.

⁸⁸³ This is the same information that CVS has refused repeatedly to provide with the G1 distributors.

distribution center anti-diversion efforts. Throughout the period, CVS allowed the controlled substance program to be severely under-resourced both in terms of total headcount and in terms of the skills and abilities of its dedicated team members (e.g., Messrs. Devlin, Mortelitti, and Miliken, etc.). Even when notified of system deficits that negatively impacted the ability of its SOM program to function, CVS failed to undertake timely and comprehensive corrective actions.

- **Tolerance of Misstatements and Errors:** Perhaps the most troubling aspect of CVS' lack of commitment, is the fact that the company allowed multiple gaps and inconsistencies in its anti-diversion program to persist unabated for periods of months or even years with no documented corrective action plan. CVS also seemed oddly comfortable allowing its employees to make obviously incorrect statements to federal regulatory authorities. This speaks to a culture where revenue generation and preventing disruption to the retail pharmacies were the valued company objectives - not compliance.

Applying the compliance maturity and program effectiveness model here, the CVS program is difficult to classify as being at the foundational level, and if the model had a remedial level, I would place the CVS program there.

12.3 Impact

The impact of CVS' failure to maintain a credible anti-diversion program is clear. During the relevant period, CVS reported no suspicious orders to the DEA regarding any of its pharmacies in Cuyahoga or Summit Counties.⁸⁸⁴ In fact the Item Review Report ("IRR") Recap Reports from January 2011 to June 2012 show that only one flagged order of hydrocodone was selected for additional due diligence in Cuyahoga and Summit Counties during that eighteen-month period.⁸⁸⁵ During this same period of time, the report shows months, where not one (i.e., zero) flagged hydrocodone order nationwide was investigated further after being listed on the IRR Report.⁸⁸⁶ In a similar vein, the IRR Recap Report for the ten-month period spanning February 6, 2013, to December 30, 2013, identified only one hydrocodone order in Cuyahoga or Summit counties that was investigated further.⁸⁸⁷

Indianapolis Distribution Center

In November 2013, the DEA notified the head of the Indianapolis distribution center ("DC"), Mark Nicastro of two stores that clearly were suspicious, but to which CVS continued shipping hydrocodone unabated. Despite this CVS claimed that the company had an effective anti-diversion and suspicious order monitoring program. To quote the DEA, these stores were:

⁸⁸⁴ See CVS RX Services, Inc.'s and CVS Indiana L.L.C.'s Objections and Responses to Plaintiffs' (First) Combined Discovery Requests to National Retail Pharmacy Defendants, Requests Nos. 3-4.

⁸⁸⁵ See IRR Recap Report, (Jan. 2011 to Jun. 2012), CVS-MDLT1-000009740.

⁸⁸⁶ See *id.*

⁸⁸⁷ See A. Burtner Deposition at 485:20-487:1; IRR Recap Report, (Feb. 6, 2013-Dec. 30, 2013), CVS-MDLT1-000010268.

CVS Store# 06880/ DEA # AH9157137 ordered [REDACTED] dosage units of Hydrocodone (Drug Code 9193) between January 1, 2012, through October 2013. Of which [REDACTED] tablets of Hydrocodone were shipped from your facility. This pharmacy is located in Vincennes, IN with a population of approximately 18,000.

Additionally, CVS Store# 6757/DEA# AH2693376 located in Columbus, IN ordered a total of [REDACTED] tablets of which your facility provided [REDACTED] tablets from January 1, 2012, through October of 2013. The population of Columbus, IN is approximately 45,000.⁸⁸⁸

12.4 Company Commitment – Compliance Culture, Organization & Resources

12.4.1 CVS' compliance record is indicative of a company that has a poor commitment to compliance.

CVS, as noted above, describes itself as “a health care innovation company helping people on their path to better health ... transforming the consumer health care experience and helping to foster healthier communities.”⁸⁸⁹ However, the company's compliance history reveals a company that has entered into a series of Corporate Integrity Agreements, 2008 (CVS Caremark),⁸⁹⁰ 2014 (CVS Caremark),⁸⁹¹ 2016 (CVS Health Corporation)⁸⁹² with the OIG for various healthcare offenses not related to controlled substances. Thus, the Justice Department has taken issue with CVS' commitment to compliance on three separate occasions.

CVS's controlled substances compliance record is also checkered. The company has settled three sets of major allegations involving controlled substances since 2009.⁸⁹³ One case involved the excessive sales of pseudoephedrine (“PSE”) by CVS stores in California and Nevada in 2007 and 2008.⁸⁹⁴ In exchange for a non-prosecution agreement and civil settlement, CVS paid \$77.6 million in civil penalties and profit forfeitures.⁸⁹⁵

⁸⁸⁸ See Email from D. Gillen to M. Nicastro, CVS Store #06880 & 6757, (Nov. 25, 2013), CVS-MDLT1-000076135.

⁸⁸⁹ See CVS HEALTH, *Investor Story*, <https://investors.cvshealth.com/investors/investor-story/default.aspx> (last visited Feb. 17, 2019).

⁸⁹⁰ See Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Caremark Corporation (Mar. 14, 2008) https://oig.hhs.gov/fraud/cia/agreements/cvs_cia_executed.pdf.

⁸⁹¹ See Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Caremark Corporation (Mar. 25, 2014), https://oig.hhs.gov/fraud/cia/agreements/ CVS_Caremark_03252014.pdf.

⁸⁹² See Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Health Corporation (Oct. 11, 2016), https://oig.hhs.gov/fraud/cia/agreements/ CVS_Health_Corporation_10112016.pdf.

⁸⁹³ CVS has settled numerous smaller cases with the DEA involving its retail pharmacy stores in Massachusetts (2015 and 2016, CVS-MDLT1-000099702 and 000060872), Texas (2014 and 2015, CVS-MDLT1-000060907 and 000060915), and Maryland (2016, CVS-MDLT1-000060805).

⁸⁹⁴ See Press Release CVS/pharmacy, *Lapse in controls of PSE sales in certain CVS/pharmacy stores in 2007 and 2008 relates to electronic monitoring system flaw that has been corrected Settlement amount fully reserved and previously disclosed; should have no further effect on company's financial results*, (Oct. 4, 2010), <https://cvshealth.com/newsroom/press-releases/cvspannounces-agreements-us-drug-enforcement-administration-and-us-attorneys-offices>.

⁸⁹⁵ *Id.*

Another case in 2013 ended when CVS agreed to pay \$11 million in civil monetary penalties to resolve recordkeeping violations involving CVS Pharmacy, Inc. and Oklahoma CVS Pharmacy, LLC.⁸⁹⁶ The DEA alleged that from October 2005 to October 2011, CVS violated a number of controlled substances recording keeping requirements including creating and maintaining “dummy” DEA registrations for prescribers and filling prescriptions for physicians lacking current or valid DEA registration numbers.

CVS, in June 2018, once more settled allegations of controlled substances violations by its pharmacies in Nassau and Suffolk counties, New York.⁸⁹⁷ Apparently, these CVS stores failed to report thefts and losses of various controlled substances including hydrocodone.⁸⁹⁸ CVS Pharmacy, Inc. agreed to pay \$1.5 million in civil penalties to resolve the matter.⁸⁹⁹

12.4.2 CVS’ penchant for complexity and compartmentalization resulted in a confusing and ineffective organizational approach to controlled substances compliance.

In general, CVS’ view of compliance simply does not enter its business or cultural equation in a meaningful way. CVS’ actions demonstrated this in three distinct, but interrelated areas:

- Its corporate entity structure,
- The assigning responsibility for controlled substances compliance, and
- Its failure to integrate its controlled substances and corporate compliance teams.

A. CVS Entity Structure

CVS Health, the corporate parent of all CVS operations, has created an extraordinarily complicated and compartmentalized entity structure. For example, CVS Pharmacy, Inc., a subsidiary of CVS Health, Inc., “owns, either directly or indirectly, all of the CVS retail pharmacies.”⁹⁰⁰ CVS also operates two separately incorporated controlled substances distribution centers relevant to this report: CVS Indiana L.L.C. located in Indianapolis, Indiana and CVS Rx Services, Inc. located in Chemung, New York. These entities were in turn owned by CVS Pharmacy, Inc.⁹⁰¹

From an operational perspective, such an entity structure creates confusion and interferes with business operations and processes that span multiple corporate entities. The entity structure does this by obfuscating

⁸⁹⁶ See Press Release, U.S. Dep’t of Justice, Drug Enforcement Admin., CVS To Pay \$11 Million To Settle Civil Penalty Claims Involving Violations Of Controlled Substances Act, (Apr. 3, 2013), <https://www.dea.gov/press-releases/2013/04/03/cvs-pay-11-million-settle-civil-penalty-claims-involving-violations-0>

⁸⁹⁷ See Press Release, U.S. Dep’t of Justice, Drug Enforcement Admin., CVS Pharmacy, Inc. to pay \$1.5 million to settle Civil Penalty Claims for violation so [sic.] the Controlled Substance Act, (Jun. 28, 2018), <https://www.dea.gov/press-releases/2018/06/28/cvs-pharmacy-inc-pay-15-million-settle-civil-penalty-claims-violation-so>.

⁸⁹⁸ *Id.*

⁸⁹⁹ *Id.*

⁹⁰⁰ See Mark Vernazza Deposition, 53:23-54:1 (Nov. 20, 2018) (Mr. Venazza is Senior Legal Counsel for CVS Pharmacy, Inc. and was the 30(b)(6) witness for the two CVS distribution centers in this case).

⁹⁰¹ See Figure 1 *infra* Appendix F.

where responsibility and authority for various operations and processes reside. As a result, corporate inertia sets in as employees do not know how to get things accomplished.

CVS further compounded these issues and challenges resulting from a compartmentalized corporate entity structure by not creating the entities that were truly separate. For example, if the Indianapolis and Chemung distribution centers were truly separate entities, then one would expect to see internal contracts or service level agreements (“SLAs”) between the distribution centers and CVS Pharmacy, Inc. setting out the distribution expectations for the individual pharmacies. Likewise, since CVS Pharmacy, Inc. owned and controlled the pharmacy ordering system, there should be SLAs between CVS Pharmacy, Inc. and the distribution centers, which depend on that system to ensure timely order fulfillment.⁹⁰² However, CVS did not have any such contracts, nor were the pharmacies required to pay for the products or the delivery services provided by Indianapolis and Chemung.⁹⁰³ Thus, ownership of the distribution process and the underlying order system were not clear cut. These are signs of poor corporate governance.

B. CVS Controlled Substances “Team”

This penchant for complexity and compartmentalization that is seen with the entity structure carried over into the organizational design of the controlled substances program. Rather than take a straight-forward approach of designating a “high-level” individual or group with sole responsibility for controlled substances compliance, CVS opted for a convoluted structure with diffuse responsibility across functional lines and corporate entities. This resulted in a controlled substances compliance “team structure” that lacked both accountability and effectiveness.

This confusion caused by CVS’ cumbersome, compartmentalized approach was readily apparent when viewed in the context of which department “owned” the SOM process. Throughout the period, “ownership” of the SOM program involved two separate functional lines within CVS: (a) Logistics and (b) Loss Prevention (“LP”). From 2006 to 2012-2013, “ownership” of the SOM program resided under Loss Prevention, which was a department located at CVS Headquarters in Woonsocket, Rhode Island.⁹⁰⁴ Sometime between 2012 and 2013, there was consensus that the Logistics Planning Department, a subset of the Logistics Department, took over “ownership” of the SOM program.⁹⁰⁵

However, there was some discrepancy as to precisely when the transfer from LP to Logistics took place. An email from Mark Nicastro in August 2013 suggested that Logistics took over well before August 2013: “My understanding is Logistics owns the process so either Dean [Vanelli] or I have it.”⁹⁰⁶ Dean Vanelli, Senior

⁹⁰² *Id.* at 48:2-6.

⁹⁰³ *Id.* at 48:17 to 52:2.

⁹⁰⁴ See Ronald Link Deposition, 19:23 to 20:1 (Dec. 11, 2018), *but see* Email from M. Nicastro to R. Link and W. Jusko, SOM Update (Aug. 18, 2013), CVS-MDLT1-000012363; Email from D. Vanelli to R. Link, *et al.*, FW 2013 Logistics Planning Initiatives (Jan. 9, 2014), CVS-MDLT1-000000459; CVS Logistics Planning Organizational Chart (Jan. 1, 2014) CVS-MDLT1-000037885.

⁹⁰⁵ See *id.* at 19:21-22; see also Dean Vanelli Deposition, 18:23-19:3; 23:7-19 (Jan. 16, 2019).

⁹⁰⁶ See Email from M. Nicastro to R. Link and W. Jusko, SOM Update (Aug. 18, 2013), CVS-MDLT1-000012363; *but cf.* R. Link, Deposition at 19:16-22 (stating the transition occurred in 2012).

Director of Logistics Planning, in a 2014 email discussing his department's 2013 initiatives stated that he assumed ownership of the SOM program.⁹⁰⁷

Similar confusion surrounded Amy Propatier's role in the SOM Program. Reference to Mrs. Propatier's SOM role can be found in the 2011 version of the CVS DEA SOP Manual, where it named her as "CVS DEA Compliance Coordinator."⁹⁰⁸ The same section also referenced Frank Devlin as the Director of Logistics Loss Prevention.⁹⁰⁹ Ronald Link, Vice President of Logistics to whom Logistics Planning reported, stated that while he did not have contact with Mrs. Propatier, he knew she was the CVS "DEA Coordinator."⁹¹⁰ According to Mrs. Propatier, "her duties" as DEA Coordinator involved filing ARCOS reports and updating the DEA SOP Manual.⁹¹¹ Consequently, as Mrs. Propatier stated, the DEA Coordinator title was "a title for reference in SOPs," and therefore was not included in her personnel records."⁹¹²

From 2006 to 2014, Mrs. Propatier's official duties were "hazardous materials specialist and logistics liaison" that, as of 2008, evolved into a "logistics Rx services manager"⁹¹³ within the Logistics Planning department. Dean Vanelli, Senior Director of Logistics Planning, claimed "[a]t no time did Amy have ownership for any DEA compliance ... with the exception of she was responsible for filing ARCOS reporting."⁹¹⁴ Mr. Vanelli's statement is supported by Logistics Planning organizational charts.⁹¹⁵

C. Integrating the Controlled Substances and Corporate Compliance Teams

Since at least 2007, there has been a Chief Compliance Officer of CVS Health, Inc., and before that CVS Caremark Corporation. From 2007 to the present, three people have occupied that post:

- Diane Nobles, Senior Vice President, Compliance & Integrity from 2007 to 2010.⁹¹⁶
- John Buckley, Senior Vice President, Chief Compliance Officer from 2011 to 2015.⁹¹⁷
- David Falkowski, current Senior Vice President and Chief Compliance Officer for CVS Health, Inc.⁹¹⁸

⁹⁰⁷ See Email from D. Vanelli to R. Link, *et al.*, FW 2013 Logistics Planning Initiatives (Jan. 9, 2014), CVS-MDLT1-000000459.

⁹⁰⁸ See CVS Distribution Center, *Controlled Drug – DEA Standard Operating Procedures Manual RX-01*, X-8, § E.3 (Nov. 8, 2011), CVS-MDLT1-000008506 ["RX-01(11/11)"].

⁹⁰⁹ *Id.*

⁹¹⁰ See R. Link Deposition at 21:5-20, 24:1-5.

⁹¹¹ See Amy Propatier Deposition 80:17 -81:2; 103:7-15; 104:5-120:21 (Nov. 29, 2018) (inquiring about any other duties she might have performed).

⁹¹² See *id.* at 79:24 to 80:2, 81:18 -82:1.

⁹¹³ See *id.* at 17:10-19.

⁹¹⁴ See D. Vanelli Deposition at 50:21-24.

⁹¹⁵ See CVS Logistics Planning Organizational Chart (Apr. 15, 2013); CVS Logistics Planning Organizational Chart (Jan. 1, 2014).

⁹¹⁶ Diane Nobles LinkedIn Profile, <https://www.linkedin.com/in/dbnobles/> (last accessed Feb. 12, 2019) (Prior to joining Walgreens, Ms. Nobles left CVS to become CCO of Walgreens from 2013-2016).

⁹¹⁷ See John Buckley LinkedIn Profile, <https://www.linkedin.com/in/jmbuckleycco/>, (last accessed Feb. 26, 2019).

⁹¹⁸ See David Falkowski LinkedIn Profile, <https://www.linkedin.com/in/david-f-45037626/> (last accessed Feb. 26, 2019).

However, no evidence was presented suggesting a linkage, much less integration, between the two compliance teams. This is even more troubling because from 2008 to the present, CVS was the subject of three separate Corporate Integrity Agreements, all mandating enhanced attention to all facets of compliance.⁹¹⁹ This, however, did not seem to include controlled substances compliance.

This failure by CVS strongly suggests that CVS focused little on controlled substances. Had CVS done so, it would have helped the anti-diversion team highlight critical issues and secure additional management support. By not doing so, CVS missed an opportunity to move towards a credible anti-diversion program. It also demonstrated a lack of company commitment to meeting its controlled substances obligations.

12.4.3 CVS also failed to properly resource its controlled substances compliance efforts throughout the period.

CVS's extraordinary fragmentation of responsibilities for the anti-diversion program also negatively impacted the resourcing for the controlled substances compliance efforts. In short, throughout the review period, the controlled substances "team" was staffed at such a low level that it was ineffective.

Starting in the 2009 timeframe, CVS centralized the review of all Inventory Review Reports ("IRRs") at a single distribution center in Lumberton, New Jersey under John Mortelliti, Loss Prevention Manager for the Lumberton Distribution Center and the mid-Atlantic region.⁹²⁰ Thus from 2009 to March 2011, CVS had only one person doing the daily review of the IRRs for the entire country.⁹²¹ As a Loss Prevention Manager, Mr. Mortelliti had no prior experience with suspicious order monitoring and did not recall any training he received.⁹²² The same was true for Frank Devlin, Director of Loss Prevention and Mr. Mortelliti's superior.⁹²³

When the finalized version of the SOM program was inserted into the DEA SOP manual in August 2010, CVS planned to have each distribution center reviewing its own IRRs.⁹²⁴ This change could have provided 11 FTEs to help handle the volume of IRR reports.⁹²⁵ However, the change was never made, and the IRR reviews

⁹¹⁹ See Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Caremark Corporation (Mar. 14, 2008) https://oig.hhs.gov/fraud/cia/agreements/cvs_cia_executed.pdf; Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Caremark Corporation (Mar. 25, 2014), https://oig.hhs.gov/fraud/cia/agreements/ CVS_Caremark_03252014.pdf; Corporate Integrity Agreement between the Office of Inspector General of the Department of Health and Human Services and CVS Health Corporation (Oct. 11, 2016), https://oig.hhs.gov/fraud/cia/agreements/ CVS_Health_Corporation_10112016.pdf.

⁹²⁰ See Henry John Mortelliti, III Deposition, 18:10-19:1 and 19:6-12, (Jan. 23, 2019).

⁹²¹ *Id.*; see also Email from J. Mortelliti to E. Demetrius RE: IRR/SOM Retunement BSR_LOG_61148 (Mar. 14, 2011), CVS-MDLT1-000057759.

⁹²² See J. Mortelliti Deposition at 19:13-22:5; 46:20-47:10.

⁹²³ See F. Devlin Deposition at 168:12-169:4.

⁹²⁴ See CVS Distribution Center, *Controlled Drug – DEA Standard Operating Procedures Manual*, VIII-7, § D.4 (Aug. 25, 2010), CVS-MDLT1-000088956 at CVS-MDLT1-000088957 ["RX-01(2010)"]; CVS Presentation, *Suspicious Order Monitoring for PSE/Control Drugs: Summary of Key Concepts and Procedures*, 7 (Aug. 27, 2010), CVS-MDLT1-000075300 ["DEA Speaking Points"];

⁹²⁵ See Email from C. Tulley to P. Hinkle, *et al.*, RE: Recap progress made, (Nov. 11, 2012), CVS-MDLT1-000055834.

continued to be done by only a few individuals at any one time.⁹²⁶ CVS claims the reason for not pushing IRR review into the individual distributions centers was “review consistency” and operational efficiency.⁹²⁷ In March 2011, the IRR review process was transferred from Lumberton to Knoxville under the control of Pam Hinkle.⁹²⁸ Upon the move to Knoxville, Ms. Hinkle ultimately gained two Loss Prevention Analysts assigned to the SOM program.⁹²⁹

The IRR workload with CVS was substantial as IRRs are generated daily for each distribution center.⁹³⁰ Therefore, in between 2009 and 2010, that meant at least 10 reports per day were generated.⁹³¹ A single sample IRR for the Indianapolis DC on November 30, 2010 contained 355 entries (71 pages x 5 lines per page).⁹³² While it is unclear whether this was a “typical” IRR,⁹³³ even if this IRR were halved, that translates into approximately 1,550 orders per day or approximately 387,500 orders per year across all distribution centers.⁹³⁴

In the summer of 2012, Aaron Burtner, SOM Manager, conducted a series of time studies to outline his average day.⁹³⁵ The results of those time studies revealed that CVS employees were doing, at best, a cursory review of the daily IRRs.⁹³⁶ Based on the number of daily orders on the IRR and Mr. Burtner’s time studies, the staffing level provided by CVS to review “flagged” orders was not sufficient to accomplish the task of conducting meaningful due diligence on orders listed on the IRR.

12.4.4 CVS’s tolerance of program documentation that was either non-existent, incomplete or inaccurate is indicative of a company with a poor compliance culture.

Throughout the review period, the CVS team operated with non-existent, incomplete or simply inaccurate program documentation. While it is not unexpected for a compliance program’s documentation to be out of sync or inconsistent with current operational practices, normally, these periods are short-lived and as the result of non-routine events, such as a merger, department reorganization, changes in personnel titles and reporting lines, etc. It also is no surprise that the frequency of these types of events correlates directly with the size and complexity of the organization. When these situations occur, the organization committed to compliance will

⁹²⁶ See Pamela Hinkle Deposition, 38:4 to 39:20 (Jan. 24, 2019)

⁹²⁷ See E mail from A. Burtner to C. Tulley and P. Hinkle, RE: Recap progress made, (Nov. 11, 2012), CVS-MDLT1-000055834; see also F. Devlin Deposition at 161:19.

⁹²⁸ See Email from J. Mortelliti to E. Demetrius, *et al.*, RE: IRR/SOM Retunement BSR_LOG_61148 (Mar. 14, 2011), CVS-MDLT1-000057759; see also P. Hinkle Deposition at 26:4-27:8; RX-01(11/11) at VIII-7, § D.4.

⁹²⁹ See P. Hinkle Deposition, 33:7-35:11.

⁹³⁰ See J. Mortelliti Deposition 40:7-9.

⁹³¹ *Id.* at 39:9-12 to 40:3-6.

⁹³² See Item Review Report, BIP006A (Nov. 30, 2010), CVS-MDLT1-000100775.

⁹³³ See J. Mortelliti at 38:22 to 39:1.

⁹³⁴ The formula is 31 pages x 5 items per page x 10 distribution centers = 1,550. Annualized the formula is 1,550 x 5 days per week x 50 weeks per year (assuming holidays and 1-week preventative maintenance shut down) = 387,500

⁹³⁵ See A. Burtner Deposition at 340:16-23.

⁹³⁶ *Id.* at 341:19 to 371:18 (including exhibits 411-418); see also Discussion *infra*.

recognize the gaps or inconsistencies and create a plan to remedy them in a timely manner (e.g., a corrective action plan).

Therefore, it is outside the norms of good corporate governance, as well as a symptom of an organization's poor compliance culture, for an organization to allow multiple gaps and inconsistencies to persist unabated for periods of months or even years with no corrective action plan. This was the case with CVS.

As previously discussed, the designation of Amy Propatier as the "CVS DEA Compliance Coordinator," an artificial position in the DEA SOP manual was an example of CVS's failure to ensure its anti-diversion program's documentation matched current practices.⁹³⁷ By providing the document naming Mrs. Propatier as the CVS DEA Compliance Coordinator, CVS misrepresented its program to a federal regulatory agency.

Likewise, in 2010 when finalizing the SOM segment of the DEA SOP manual, CVS developed and formally approved a set of speaking points "for the DEA agents if they come ... and question suspicious monitoring."⁹³⁸ Although acknowledging that the slide deck could be shared with the DEA, Mr. Mortelliti, nevertheless, cautioned: "Please be sure your team understands it before presenting, **so it doesn't look like a prop instead of a tool.**"⁹³⁹

The DEA Speaking Points slide deck stated "DC Rx – Review Report (IRR) Daily and determine whether variances are within acceptable ranges."⁹⁴⁰ According to Frank Devlin, "DC Rx" is a reference to the distribution center pharmacy.⁹⁴¹ However, the DEA SOP manual issued two days before the date of the DEA Speaking Points slide deck stated: "Currently the Item Review Report (IRR) for control drugs is being reviewed at a central location in NJ. During the month of September 2010, the report will be transitioned to each pharmacy DC ..."⁹⁴² This never occurred, and the DEA SOP manual in November 2011 reflected this situation stating "Currently the Item Review Report (IRR) for control drugs is being reviewed at a central location in Knoxville, TN."⁹⁴³ Thus, the DEA Speaking Points misrepresented the CVS SOM program when it was created and approved to be given to the DEA.

⁹³⁷ See Discussion and notes *infra*

⁹³⁸ See Email from J. Mortelliti to M. Jamagin, FW: DEA Speaking Points (Sept. 1, 2010), CVS-MDLT1-000075299.

⁹³⁹ *Id.* (emphasis added).

⁹⁴⁰ See DEA Speaking Points at 7.

⁹⁴¹ See F. Devlin Deposition at 139:2-5

⁹⁴² See RX-01(2010) at VIII-7, § D.4.

⁹⁴³ See RX-01(11/11) at VIII-7, § D.4.

12.5 Program Core – Requirements, Education, Detection & Corrections

12.5.1 Overall CVS, from 2006 to 2014, maintained an incomplete and dysfunctional anti-diversion program related to its distribution of controlled substances.

Although CVS had an advantage over other controlled substances distributors in that the “customers” of its distribution centers were CVS-owned or operated pharmacies, CVS failed to design, implement, operate or maintain an effective anti-diversion program that included the detection and reporting of suspicious orders. CVS and its distribution centers were in the enviable position that access to pharmacy dispensing data was in-house and accessible. However, rather than capitalize on this differentiator by creating an effective program, CVS chose to do as little as possible with regards to controlled substances compliance.

A. 2006 – 2009

During the period from 2006 to 2009, CVS’ anti-diversion program hinged on two main processes: “Pickers and Packers” and PDMR reports. However, neither process adequately met the basic parameters of a credible SOM system.

1. Pickers and Packers

The pickers and packers (“P&Ps”) were distribution center employees, “who actually pick the drugs, place them in secured totes and see to it that those, then, are transferred for loading on trucks.”⁹⁴⁴ With the absence of documented criteria for determining when an order was suspicious, the P&Ps were forced to rely on their “gut feeling” or other unvalidated informal rules.⁹⁴⁵ For example, Ellen Wilson, who worked in one of the Rx department picking aisles at the Indianapolis DC, described using a rule taught to her by Charlotte Rucker based on hydrocodone bottle sizes:

She said, [a]s a rule you send out 12 -- no more than 12 of the little ones, six of the big ones and two to three of the bigger -- the large size ones as a rule across the board.⁹⁴⁶

Ms. Wilson was not aware of how many different formulations and strengths that hydrocodone came in and therefore to her, it was permissible to “order 11 little bottles of hydro all different doses and [that] would be

⁹⁴⁴ See M. Vernazza Deposition at 165:16-20.

⁹⁴⁵ See E. Wilson Deposition 17:11 - 18:20, 62:3-19; 63:19-24 (January 24, 2019); see also G. Milikan Deposition, 147:9-15 and 149:18 -150:3. Training for pickers and packers was only seen referenced in documents dating from August 2013.⁹⁴⁵ The “huddle” as the training was called, was intended to remind the pickers and packers about the duty to monitor orders of controlled substances. See Email from A. Propatier to J. Meikle, *et al.*, SOM Work Instructions (Aug. 6, 2013), CVS-MDLT1-000003020; Work Instructions for Suspicious Order Monitoring, CVS-MDLT1-000003020 at CVS-MDLT1-000003021 and DC Huddle, CVS-MDLT1-00003028. However, beyond a general reminder, the “huddle” document does not list specific criteria that should be used when judging whether an order is deemed suspicious or not. *Id.*

⁹⁴⁶ See E. Wilson Deposition at 63:19-24.

okay ...⁹⁴⁷ There was also no consideration of what previously was shipped to the pharmacy at all.⁹⁴⁸ As a result very few orders were ever flagged by the P&Ps.⁹⁴⁹

The CVS system of using P&Ps as a suspicious order monitoring control is completely ineffective as the P&Ps had no visibility to order histories to evaluate discrepancies in size, frequency or pattern, the fundamental criteria established by the DEA for suspicious orders.⁹⁵⁰

2. PDMR Reports

The PDMR reports (a.k.a. a VIPER reports) were loss prevention reports that “reflected orders in the aggregate and compared those orders to dispensing and looked to see if there were increases in ordering over dispensing, coupled with other indicia that may prompt loss prevention personnel to undertake an investigation.”⁹⁵¹ Therefore, they were reports to show how much was shipped to and dispensed from a pharmacy and whether there was a theft of product.⁹⁵² There was no evidence presented that formal documented standards governing the use of PDMR reports ever existed.⁹⁵³

While CVS represented that PDMR reports were part of the company’s efforts to monitor suspicious orders, Mark Vernazza testified:

[T]his report was not what we deemed a suspicious order monitoring report. It’s relevant to orders and order size and, some degree, order pattern. But the point of this was not to produce results for the purposes of determining whether suspicious orders were made and reporting those to the DEA.⁹⁵⁴

Furthermore, the reports were deficient as a SOM reports in two important respects. First, the reports did not provide data on unusual order size, frequency or pattern.⁹⁵⁵ Second, they were produced at monthly intervals or more and thus provided only a retrospective view that could not be used to block and hold suspicious orders.⁹⁵⁶ Therefore, the PDMR reports also were an ineffective anti-diversion control when measured against the DEA’s requirements to prevent contemporaneous diversion.

⁹⁴⁷ *Id.* at 67:16-21.

⁹⁴⁸ *Id.* at 71:17- 72:13.

⁹⁴⁹ *See* Sherri Hinkle Deposition, 83:23 to 86:5 (Jan. 25, 2019) (perhaps one order every six months was flagged).

⁹⁵⁰ *See* 21 C.F.R. § 1301.74(b).

⁹⁵¹ *See* M. Vernazza Deposition at 178:20 to 179:4.

⁹⁵² *See* Aaron Burtner Deposition at 384:12-22.

⁹⁵³ *See* F. Devlin Deposition at 420:13-23.

⁹⁵⁴ *See* M. Vernazza Deposition at 191:14-21.

⁹⁵⁵ *See* A. Burtner Deposition at 383:10 to 384:9.

⁹⁵⁶ *See* M. Vernazza Deposition at 192:2-12.

B. 2009 - 2011

The period of 2009 to 2011 saw perhaps the most substantive changes to the CVS SOM and anti-diversion programs. These changes included written standards, automated threshold detection systems, and standardized reporting of “flagged” orders. Despite these modifications, CVS’ convoluted implementation of these changes still resulted in an ineffective system to detect, stop and report suspicious orders

1. Controlled Drug – DEA SOP Manual

In early 2007, CVS, with the help of the BuzzeoPDMA group, began work on an SOP manual that was intended to cover all facets of DEA controlled substances compliance, including suspicious order monitoring.⁹⁵⁷ However, by November, neither the final manual nor the SOM section was complete “[w]e are still in the process of writing the Suspicious Order Monitoring Section of the SOP.”⁹⁵⁸ Later, when the first version of the SOP manual finally was issued in December 2007, the SOM section still remained incomplete.⁹⁵⁹

By 2009, almost two years after the SOP manual was first distributed, CVS still had not completed the SOM section: “the SOM section is still not incorporated in the SOP, in the event of an audit and the question comes up please direct them to corporate ... for the explanation of the program.”⁹⁶⁰ As Mr. Mortelliti wrote in November 2009, “I am trying to get a rough draft SOM SOP to you prior to the meeting.” and he recognized that “[t]his is a big issue with CVS and the DEA.”⁹⁶¹ However, despite recognizing this was a big issue, CVS did not incorporate the final missing section until the end of August 2010, and then did so only because of a need to respond to an apparent promise to provide it to the DEA.⁹⁶²

Adding further complexity and confusion to CVS’ written standards was the fact that the DEA SOP Manual was revised at least four more times between August 2010 and November 2011.⁹⁶³ So many revisions in such a short time frame is indicative of a compliance program that exhibits poor self-reflection and weak critical thinking. Being in an almost constant state of flux, made it extremely difficult for CVS to ensure the manual’s recipients were working with the most current version of the document as well as have all applicable employees trained.

⁹⁵⁷ See Letter G. Glatz to F. Devlin, Regulatory Consulting Services, 2 (Mar. 22, 2007) (At that time BuzzeoPDMA was a division of Dendrite, which was later acquired by Cedegim), CVS-MDLT1-000109199 [“BuzzeoPDMA Proposal”]; see also Email from A.L. Brown to A. Brumfield, *et al.*, New RX DEA SOP (Nov. 27, 2007) (referring to the *Controlled Drug – DEA Standard Operating Procedures Manual RX-01*), CVS-MDLT1-000025204; CVS Distribution Centers, *Controlled Drug – DEA Standard Operating Procedures (SOPs) Manual*, Cover page (Nov. 2007) (Showing seven drafts from May 1, 2007 to November 2007), CVS-MDLT1-000025204 at CVS-MDLT1-000025206 [“2007 DEA SOP”].

⁹⁵⁸ See also Email from A.L. Brown to A. Brumfield, *et al.*, New RX DEA SOP (Nov. 27, 2007), CVS-MDLT1-000025204.

⁹⁵⁹ See RX-01(2009).

⁹⁶⁰ See Email from A. Propatier to W. McDaniels, *et al.*, Updated DEA SOP (Apr. 3, 2009) (A. Propatier was formerly A.L. Brown), CVS-MDLT1-000066574.

⁹⁶¹ See Email from J. Mortelliti to C. Knight, RE: November 10, 2009 (Nov. 5, 2009), CVS-MDLT1-000087889.

⁹⁶² See RX-01(2010); see also Email from A. Propatier to A. Lamoureux, DEA SOP 08-25-10.doc (Aug. 26, 2010), CVS-MDLT1-000088956; Email from J. Mortelliti to F. Devlin, *et al.*, RE: DEA SOP (Aug. 23, 2010) (referencing Mr. Devlin’s earlier email to him stating “we promised this to DEA by Wednesday.”), CVS-MDLT1-000089188.

⁹⁶³ See RX-01(11/11) at 1, CVS-MDLT1-000008506 (showing revisions for 8/25/10, 11/29/10, 3/11/11, 5/6/11 and 11/8/11).

Also, the fact that CVS revised this manual so frequently (in some cases three or more times per year) suggests that the company could not draft the SOP closely enough to mirror actual operations, or alternatively that CVS kept “tweaking” and testing the process in the hope they could obtain preordained outcomes. Based on how CVS handled the Buzzeo algorithm (discussed later), I believe that the latter is the more plausible explanation.

When it was finally issued in August 2010, the newly revised SOM section covered several topics, including (a) items reviewed, (b) product buildup, (c) the Item Review Report or IRR, (d) review escalation steps, and (e) the suspicious order reports.⁹⁶⁴ It also expressly mandated that all CVS distribution centers (“DCs”) “must follow these procedures.”⁹⁶⁵

In the Items Reviewed section, CVS maintained that:

CVS has established Control Drug order thresholds which will flag on the IRR (Item Review Report) These thresholds are **the primary tool** to prevent stores from purchasing excessive or potentially suspicious Control Drug orders. These thresholds are based on historical trends of sales. Stores may order more than the historical average; however, the DC may not ship amounts that exceed these thresholds if it is believed to be suspicious.⁹⁶⁶

Adopting the prior version of the manual’s use of vague language, the section only outlined two possible factors to be considered for determining when an order “is believed to be suspicious,” although the DC Pharmacy Supervisor may consider more: (a) known reasons for increased orders such as product shortages or (b) “Corporate promotional activities.”⁹⁶⁷ Citing just these two examples was disingenuous given the DEA’s diversion factors were sent to all registrants in September 2006 and easily could have been incorporated in this 2010 SOP version.⁹⁶⁸

2. Cegedim Compliance Solutions Suspicious Order Monitoring System (“CCS-SOMS”)

In March 2007, BuzzeoPDMA proposed and ultimately was engaged for a series of projects with CVS to help with developing an enhanced anti-diversion program focused around suspicious order monitoring.⁹⁶⁹ As part of the overall project plan, BuzzeoPDMA planned to “develop a specific significant loss threshold methodology and prepare an SOP specific to controlled substances managed and distributed by CVS” by utilizing “historical data from CVS’s distribution facilities in developing the methodology and the schedule specific significant loss thresholds.”⁹⁷⁰ This work was part of a larger project to assist “CVS in the development of a comprehensive

⁹⁶⁴ See RX-01(2010) at VIII-6 to VIII-8, §§ D.1-7 (Control Drug Suspicious Orders).

⁹⁶⁵ *Id.* at VIII-6, § D.1.

⁹⁶⁶ *Id.* at VIII-7, § D.2.

⁹⁶⁷ *Id.* at VIII-7, § D.4(a-b).

⁹⁶⁸ See DEA 9/27/2006 Letter at 2, CVS-MDLT1-000010552.

⁹⁶⁹ See generally BuzzeoPDMA Proposal.

⁹⁷⁰ *Id.* at 2.

suspicious order monitoring and compliance plan that will allow CVS to operate a system to disclose suspicious orders of controlled substances,” with an automated computer program at its core.⁹⁷¹

With the purchases of BuzzeoPDMA by Cegedim, the automated program became known as the Cegedim Compliance Solutions Suspicious Order Monitoring System or CCS-SOMS. The CCS-SOMS program was “designed to evaluate orders and determine whether they are more likely to fit the DEA's definition of a ‘suspicious order’ or less likely to fit the DEA's definition of a ‘suspicious order.’”⁹⁷² If an order is more likely to fit the DEA’s definition of a “suspicious order,” the CCS-SOMS program “pends” or flags an order that may be suspicious.⁹⁷³ This determination is made by scoring the order.

The scoring methodology or algorithm was the heart of CCS-SOMS program. The order’s score was the result of scoring each order line item against a series of attributes (e.g., order qualities).⁹⁷⁴ These attributes included “markers or data calculated from a twelve-month historical database [and] ... identifiers - binary variables that must be either yes (assigned a value of 1) or no (assigned a value of 0).”⁹⁷⁵ Certain factors had weighted values (e.g., a coefficient) if they were indicative of a suspicious order.⁹⁷⁶ The CCS-SOMS program sought “to apply statistical techniques to establish ‘norms’ and ‘deviations’ in order that the overall ‘suspiciousness’ of the order [could] be evaluated,” therefore “[a]t its core, the system [used] a heavily modified logistic regression model.”⁹⁷⁷ In addition, “[t]he model [was] designed so that any order with a score of 0.15 or higher is identified as suspicious, pending, and should be investigated further.”⁹⁷⁸ Cegedim delivered the first version of the CCS-SOMS system to CVS in December 2008.⁹⁷⁹

3. Item Review Reports (“IRRs”)

With the implementation of the CCS-SOM system, the IRR became the vehicle by which “pending” orders, (i.e., orders that scored above 0.15 and thus were “suspicious”) were reported out of the system to Loss Prevention. According to the DEA SOP Manual, the IRR report was “an analysis of all Control Drug orders from the stores within the prior 24 hours ... [that] identifies orders that are statistically significant or that vary from historical monthly trends based on the previous 6 months as well as the current month.”⁹⁸⁰

⁹⁷¹ *Id.* at 3.

⁹⁷² See Compliance Solutions Powered by BuzzeoPDMA, *Descriptive Overview Document Cegedim Compliance Solutions Suspicious Order Monitoring Controlled Substances “Retunement”*, 2 (Feb. 2011) (email attachment from H.J. Mortelliti to F. Devlin, FW: The CVS Retunement, (Feb. 2, 2011)), CVS-MDLT1-000114642 [“Retunement Document”].

⁹⁷³ *Id.*

⁹⁷⁴ *Id.*

⁹⁷⁵ *Id.*

⁹⁷⁶ *Id.*

⁹⁷⁷ *Id.* at 3.

⁹⁷⁸ *Id.* at 3 (emphasis in original removed).

⁹⁷⁹ *Id.* at 1.

⁹⁸⁰ See RX-01(2010) at VIII-7, § D.4.

The SOP manual also mandated that “The DC Pharmacy Supervisor has primary responsibility for reviewing the report and investigating all orders that may be excessive or unusual.”⁹⁸¹ However, despite how use of the IRR was portrayed in the SOP, CVS simply did not follow its own SOP.

As set out in the SOP, the plan to decentralize SOM responsibilities to the individual distribution centers was never implemented.⁹⁸² Although the IRR review process was centralized first in the Lumberton DC, it was later transferred to the Knoxville DC and subsequently centralized in Indianapolis.⁹⁸³ The reasoning behind not decentralizing the SOM process will be discussed later in this report, but its negative impact on the SOM program cannot be overemphasized.

IRR reports were generated daily for each distribution center.⁹⁸⁴ According to Mr. Mortelliti, who was reviewing all the daily IRR reports, it was his practice to “freeze” every flagged hydrocodone order that appeared on the IRR by contacting the Loss Prevention Manager and the Pharmacy Manager at the relevant Distribution Center.⁹⁸⁵ He stated his process was to also contact the Field Viper Analyst (“FVA”) or the Regional Loss Prevention Manager (“RLPM”) in order for them to conduct an investigation.⁹⁸⁶

Despite Mr. Mortelliti’s contention that CVS froze and investigated every hydrocodone order listed on the IRRs, other testimony and evidence reviewed does not support his claims. At the outset, CVS did not produce, nor could Mr. Mortelliti cite to, any copies of IRRs or other documentation showing that he sent flagged orders to the FVAs for follow up.⁹⁸⁷ In fact, Terrence Duggar, the Loss Prevention Manager for the Indianapolis DC,⁹⁸⁸ recounting a conversation with a DEA agent during an August 2010 site inspection, stated in a contemporaneous email that:

I shared with her [the DEA agent] the Suspicious Order Monitoring report (IRR) and she asked how often I received it. I told her daily and weekly, but I have not received the file in a few months as the report was being tweaked. I told her that it was monitored corporately by John Mortelliti. She asked what happens when he calls regarding information on the report, I told her that **I have never received a call regarding information from the report.**⁹⁸⁹

Mr. Duggar also testified that he never monitored any controlled substances.⁹⁹⁰

⁹⁸¹ *Id.*

⁹⁸² See RX-01 (2010) at VIII-7 at § 4 (“During the month of September 2010 the report will be transitioned to each pharmacy DC ...”); Email from J. Mortelliti to T. Janson, *et al.*, RE: VBDC question (Oct. 12, 2010) (“I will be doing the control drug IRR for the network so there won’t be as much confusion trying to decipher the report in its current form.”), CVS-MDLT1-000075542.

⁹⁸³ See Discussion *infra*.

⁹⁸⁴ See J. Mortelliti Deposition 40:7-9.

⁹⁸⁵ See J. Mortelliti Deposition at 57:16-58:12.

⁹⁸⁶ See J. Mortelliti Deposition at 54:4-13; 55:2-10; 56:18-24.

⁹⁸⁷ See J. Mortelliti Deposition at 302:9 to 303:14.

⁹⁸⁸ See Terrence Duggar Deposition at 20:15-22:8 (Jan. 23, 2019) (Mr. Duggar was the LP Manager from 2005 to November 2010).

⁹⁸⁹ See Email from T. Duggar to F. Devlin, *et al.*, DEA day 3 (Aug. 26, 2010) (emphasis added), CVS-MDLT1-000010223; *see also* T. Duggar Deposition at 114:10-20.

⁹⁹⁰ See T. Duggar Deposition at 22:10-15.

With regards to investigating orders, the evidence suggests that CVS in practice rarely investigated flagged orders. This can be seen in the IRR Recap Report. The IRR Recap Report was a periodic report developed by CVS that collected all orders flagged by the daily IRRs over a given time period that were subjected to additional investigation (a.k.a. due diligence).⁹⁹¹

The report for January 2011 through June 2012 shows that Mr. Mortelliti deemed very few hydrocodone orders as needing additional investigation.⁹⁹² Furthermore, it is unclear exactly what type of investigation was performed by the FVAs and RLPMs as Mr. Mortelliti did not maintain any record of their investigations except “when they told [him] it was okay to release the order.”⁹⁹³ He also stated that he thought they used the PDMR (VIPER) reports in their investigation.⁹⁹⁴ However, there were no policies and procedures governing how these investigations were conducted to corroborate exactly what was done.⁹⁹⁵

By not following the IRR process as described in the DEA SOP Manual, CVS failed to hold, investigate and report suspicious orders identified by the CCS-SOMs system and recorded on the IRRs. CVS failed to recognize that any orders identified by the CCS-SOMs system and placed on the IRR report were by definition “suspicious” and needed further investigation before being shipped to the pharmacy. In addition, CVS’ “token” due diligence did not constitute a credible investigation. Thus, the process as implemented by CVS did not meet the requirements and expectations for an adequate suspicious order monitoring program as defined by the DEA.

4. “Tweaking” the CCS-SOMs System

Faced with the mounting problem that the new system was flagging large numbers of orders, whether real false positives or not, the CVS solution was to “tweak” the CCS-SOMs system to flag orders at a manageable level. In other words, rather than increase headcount to support the SOM program or alternatively taking an in-depth look at its pharmacies’ ordering practices to determine what might be the root cause of the problem, CVS simply altered the system to force the desired outcome even though doing so compromised the effectiveness of the CCS-SOMS program.

As recounted by the Cegedim team:

In July 2009, CVS Staff advised that the current SOM model was “pending” a large number of orders that were **not suspicious on their face** and were ‘cleared’ by [the] CVS staff. This can infrequently occur when the model uses data from a fixed, unchanging period of time prior to the model’s initial deployment. In light of CVS’ **perceived number of “false positives,”** CCS

⁹⁹¹ See, e.g., A. Burtner Exh. 440, CVS-MDLT1-000010268 (2013); IRR Recap Report, (Jan. 2011 to Jun. 2012), CVS-MDLT1-000009740.

⁹⁹² See IRR Recap Report, (Jan. 2011 to Jun. 2012), CVS-MDLT1-000009740. The Recap Report shows flagged orders for control drugs (e.g., hydrocodone) as well as pseudoephedrine (“PSE”).

⁹⁹³ See J. Mortelliti Deposition at 92:8-9.

⁹⁹⁴ See *id.* at 67:13-68:1.

⁹⁹⁵ See *id.* at 199:8 to 200:13; see also RX01(11/11) at VIII-7 to VIII-8 § D.5.

statisticians made revision to the CVS model through an adjustment to the algorithm “coefficients.”⁹⁹⁶

The coefficient adjustment was provided to CVS in late August 27, 2009.⁹⁹⁷ However, by February 2010, CVS determined that the coefficient revisions did not go far enough, and the Cegedim team suggested raising the magnitude of the score from 0.15 to something higher “in small increments.”⁹⁹⁸ Cegedim also counseled CVS that it was “important to document what you are doing in a way to show the DEA or other authorities that the changes to the pended orders make sense and that the new policy/procedures are based upon too many unwarranted investigations.”⁹⁹⁹

Despite Cegedim’s cautions, CVS proceeded to aggressively test higher magnitudes, up to 0.21 at first,¹⁰⁰⁰ but the IRR was “still large even for the most aggressive formula.”¹⁰⁰¹ CVS finally settled on 0.65 in July 2010, in part because Mr. Mortelliti concluded that he “could find **not one** item worthy of investigation below .65.”¹⁰⁰² However, it appears that Mr. Mortelliti still was not completely satisfied as he suggested that 0.70 “look[ed] a bit more realistic.”¹⁰⁰³

Cegedim, however, expressed its concern writing to CVS stating “[t]hat’s quite a departure from the initial threshold, “and suggested that CVS should undertake a “retunement” of the system.”¹⁰⁰⁴ Even after the February 2011 retunement, which reset algorithm for flagging orders back to the 0.15 level, CVS appears quickly to have returned to operating at the 0.65 level.¹⁰⁰⁵

5. Lost Order Data

In October 2010 yet another problem with the system was discovered, which ended CVS’ attempt to decentralize the IRR review process.¹⁰⁰⁶ As Mr. Mortelliti described it:

DEA expects CVS to prevent suspicious orders from being filled out of our DC’s. The current IRR does not provide the proper information to meet the DEA’s needs. We need control drugs to

⁹⁹⁶ See Retunement Document at 1 (emphasis added).

⁹⁹⁷ See Email from F. Devlin to E. Demetrius, FW: SOM Report (Aug. 27, 2019), CVS-MDLT1-000109623; *see also* Retunement Document..

⁹⁹⁸ See Email from R. Williamson to F. Devlin, Adjustment to CVS SOM (Feb. 15, 2010), CVS-MDLT1-000110439 at CVS-MDLT1-000110441.

⁹⁹⁹ *Id.*

¹⁰⁰⁰ See Email from A. Santhoshraj to J. Mortelliti, RE: Adjustment to the CVS SOM (Mar. 10, 2010), CVS-MDLT1-000110439.

¹⁰⁰¹ See Email from J. Mortelliti to F. Devlin, FW: Adjustment to the CVS SOM (Mar. 5, 2010), CVS-MDLT1-000111260.

¹⁰⁰² See Email J. Mortelliti to R. Williamson, RE: SOM Update (Jul. 26, 2010) (emphasis added), CVS-MDLT1-000088734.

¹⁰⁰³ See *id.* at CVS-MDLT1-000088735.

¹⁰⁰⁴ See Email from R. Williamson to J. Mortelliti, *et al.*, RE: SOM Update (Jul. 26, 2010), CVS-MDLT1-000088734 at CVS-MDLT1-000088735.

¹⁰⁰⁵ See J. Mortelliti Deposition at 327:12 to 330-24.

¹⁰⁰⁶ See Email from J. Mortelliti to T. Janson, *et al.*, RE: VBDC question (Oct. 12, 2010) (“I will be doing the control drug IRR for the network so there won’t be as much confusion trying to decipher the report in its current form.”), CVS-MDLT1-000075542.

be monitored by “active ingredient.” Currently the control drugs are monitored by item. The IRR loses all order history when the info on the item changes **causing CVS to be non compliant [sic.] with DEA expectations.**¹⁰⁰⁷

The CCS-SOMs system was dependent upon historical data for accuracy, and as Cegedim told CVS “[t]he “retunement” event is a recommended practice to review and possibly re-adjust the SOM model coefficients ... **since the model is developed using historical data** that is provided at the start of the design and ordering habits may naturally evolve and change over time.”¹⁰⁰⁸

By capturing data by item versus active ingredient, the system was vulnerable to changes in the product information such as minor name changes:

We thought this would be a great idea at the time but what we found was that the system cannot match historical data to an item if the manufacturer changes the name of the item Example, Hydro 5 mg can be changed to Hydro mg5. Same item just put the 5 in front of mg. The system cannot match this item because of the change and therefore loses historical data.¹⁰⁰⁹

Despite this gap being identified as a high priority with regulatory compliance implications, CVS failed to remedy the problem in a timely manner.¹⁰¹⁰ More than seven months (April 2011) after Mr. Mortelliti’s “business idea” was submitted, the CVS IT Logistics Team was still inquiring how to prioritize this critical project with the 58 other projects in the Team’s work queue¹⁰¹¹ despite a plea from Mr. Mortelliti to the IT Team and to his supervisor, Mr. Devlin, in October 2010 to expedite the fix.¹⁰¹²

During this prolonged gap period, Mr. Mortelliti reviewed “the Control Drug IRR [based] on **commonsense** as apposed [sic.] to **IRR Historical Data.**” and by trying to manually retrieve historic order data from prior IRRs.¹⁰¹³ As Mr. Mortelliti noted, “I know, that this is scary ...” because the “gap” reduced the effectiveness of the CCS-SOMs even further beyond the high magnitude score as the lack of the historical data rendered the algorithmic score for an order unreliable.¹⁰¹⁴

¹⁰⁰⁷ See John Mortelliti, *Control Drug IRR update*, CVS Pharmacies Business Idea Description (Oct. 6, 2010; MetaData for last revision) (emphasis added), CVS-MDLT1-000034175 [“Business Idea Description”]. In Mr. Mortelliti’s deposition the document was identified as being dated October 8, 2010. See J. Mortelliti Deposition, 129:11-12.

¹⁰⁰⁸ See Retunement Document at 2 (emphasis added).

¹⁰⁰⁹ See Email from J. Mortelliti to T. Janson, *et al.*, RE: VBDC question (Oct. 12, 2010), CVS-MDLT1-000075542.

¹⁰¹⁰ See Email from G. Misiaszek to J. Andrade, RE: Logistics Business Support Requests 2011–04-28 (Apr. 29, 2011); CVS-MDLT1-000029864. The attached project plan chart shows a projected completion date of December 31st, 2011).

¹⁰¹¹ See Email from G. Misiaszek to J. Andrade, RE: Logistics Business Support Requests 2011–04-28 (Apr. 29, 2011); CVS-MDLT1-000029864.

¹⁰¹² See Email from J. Mortelliti to G. Misiaszek, *et al.*, Control Drug IRR important info, (Oct. 6, 2010) (requesting help to expedite the fix), CVS-MDLT1-000034168 at CVS-MDLT1-000034169.

¹⁰¹³ See *id.*; see also J. Mortelliti Deposition at 148:13-149:3 (noting he was alarmed at the amount of effort that was necessary to get the historical data).

¹⁰¹⁴ See J. Mortelliti Deposition at 160:6-162:4.

C. 2011 to 2014

From 2011 to 2013, CVS aggressively began to make changes to its anti-diversion program and SOM process. However, despite the additional efforts, these changes overall failed to improve the program in a meaningful way. Once more, CVS' convoluted implementation of these improvements simply achieved an ineffective system to detect, report and stop suspicious orders which is neither good compliance nor in conformance with the DEA's expectations.

1. IRR Review Shift to Knoxville

By March 2011, although CVS continued to maintain a centralized IRR review process, responsibility shifted from Mr. Mortelliti at Lumberton DC to Ms. Hinkle's team in the Knoxville DC.¹⁰¹⁵ Initially Shannon Miller, Loss Prevention Supervisor was tasked with the daily IRR reviews, but ultimately, Ms. Hinkle gained two Loss Prevention Analysts assigned to the SOM program.¹⁰¹⁶ As discussed previously, although Ms Hinkle had these additional resources to support the SOM program, the program remained severely under-resourced.¹⁰¹⁷

2. Use of MicroStrategy

Although it is not completely clear, it appears that the any due diligence follow-up on suspicious orders in the IRR was extremely limited and the only tools available for the due diligence follow-up prior to 2012 were the PDMR (VIPER) reports.¹⁰¹⁸ Beginning in February 2012, CVS began incorporating a program called MicroStrategy into the SOM process.¹⁰¹⁹

According to Aaron Burtner, former CVS SOM Manager, MicroStrategy became the primary tool used to investigate flagged orders.¹⁰²⁰ Analysts using MicroStrategy had access to information on patient ID number, doctors, how the drugs were paid for (e.g., cash or insurance), dispensing data, the patient population that was purchasing the drug, information on the pharmacy, and information on the patient.¹⁰²¹ Thus, the MicroStrategy tool was a significant improvement for order investigation. If MicroStrategy was used, the process was to make note of it and attach it to the IRR report.¹⁰²² The presumption was that if there were no notes attached to the IRR, the analyst did not use MicroStrategy.¹⁰²³

¹⁰¹⁵ Email from J. Mortelliti to E. Demetrius, *et al.*, RE: IRR/SOM Retunement BSR_LOG_61148 (Mar. 14, 2011), CVS-MDLT1-000057759; *see also* P. Hinkle Deposition at 26:4 to 27:8; RX-01(11/11) at VIII-7, § D.4.

¹⁰¹⁶ *See* P. Hinkle Deposition, 38:4-39:20.

¹⁰¹⁷ *See* Discussion *infra*.

¹⁰¹⁸ *See* J. Mortelliti Deposition at 67:13-68:1.

¹⁰¹⁹ *See* CVS, IRR SOM Review Project Plan Phase 1, ID 17 and 21 (undated) (showing that use of MicroStrategy started on or about 2/28/12), CVS-MDLT1-000114321.

¹⁰²⁰ *See* A. Burtner Deposition at 375:4-9.

¹⁰²¹ *See* A. Burtner Deposition at 396:23-397:17; *see also* Shauna Helfrich Deposition, 176:3-12 (Jan. 10, 2019)

¹⁰²² *See* A. Burtner Deposition at 308:22-309:2; S. Helfrich Deposition at 25:17-21.

¹⁰²³ *See* A. Burtner Deposition at 309:4-9.

CVS, however, limited the use of MicroStrategy to those cases where the analysts conducted additional due diligence or a “deep-dive” on a particular IRR order.¹⁰²⁴ “Deep-dives” were not performed on every order flagged on the IRR, but only on a subset of orders chosen by the daily IRR reviewer.¹⁰²⁵

According to Aaron Burtner, his review of the IRR report was merely “double-checking the algorithm,” rather than a true review based on additional data.¹⁰²⁶ As a result, the selection of orders for additional due diligence was in essence random. Since the use of MicroStrategy was limited, CVS simply continued its practice of poor due diligence.

3. AGI's Store Metrics

By October 2012, CVS was not satisfied with the outcome of Cegedim's efforts to address the problems with the CCS-SOM system and so brought AGI on board to “develop an algorithm to fix issues with [the] existing algorithm used for the SOM system.”¹⁰²⁷ However, as CVS informed AGI, the algorithm was not the only issue needing attention. During the initial discussion and planning phase, CVS told AGI that:

1. “For consistency, the SOM process is reviewed at a central location for all 11 RX DC's.”
2. CVS was using eight (8) algorithms.
3. Out-of-stock products could be ordered from an outside vendor, but those orders “are not pushed through the SOM process.”
4. The IRR process identified “irregular orders,” which were subject to further (unspecified) review, and the DC was contacted not to ship the order until cleared.¹⁰²⁸

Thus, even as late as October 2012, CVS's anti-diversion efforts continued to be plagued by severe deficiencies undermining any efforts to achieve a good controlled substances compliance program.

Delivered at the end of 2012, AGI Store Metrics was the successor program to the MicroStrategy tool.¹⁰²⁹ While the Store Metrics program provided the same information as MicroStrategy, its primary advantage was that it put that information into one combined dashboard for review¹⁰³⁰ and “greatly reduced the amount of time required to review these [flagged] orders as the [Store Metrics] report generates in a few seconds rather than a few minutes required for MicroStrategy.”¹⁰³¹ CVS replicated the MicroStrategy process, and analysts using

¹⁰²⁴ See A. Burtner Deposition at 267:5 to 268:16, 308:16-20.

¹⁰²⁵ See A. Burtner Deposition at 268:17 to 269:1.

¹⁰²⁶ See A. Burtner Deposition at 459:19 to 460:15.

¹⁰²⁷ See Email from P. Hinkle to F. Devlin, Conference Call Notes - 10 5 12.docx, (Oct. 5, 2012) (referencing notes taken by Aaron Burtner, SOM Manager), CVS-MDLT1-000033579.

¹⁰²⁸ See AGI/ CVS Discussion, Pharmacy/DC Ordering Process conference call recap, 1-2 (October 5, 2012), CVS-MDLT1-000033580.

¹⁰²⁹ See Email from A. Burtner to D. Vanelli and P. Hinkle, Attorney Client Privilege – Store Metrics Report, (Dec. 20, 2012), CVS-MDLT1-000109775; see also A. Burtner Deposition at 394:1-5 (Store Metrics becoming the go-to report).

¹⁰³⁰ See A. Burtner Deposition at 311:14-18.

¹⁰³¹ See Email from A. Burtner to D. Vanelli and P. Hinkle, Attorney Client Privilege – Store Metrics Report, (Dec. 20, 2012), CVS-MDLT1-000109775.

Store Metrics would attach the reports to the IRR, and if an IRR had no Store Metrics report attached, it was an indication that the analyst had not used Store Metrics.¹⁰³² However, the new program was not without its own problems.

As Mr. Burtner communicated to Mr. Vanelli and Ms. Hinkle in late 2012, “the dispensing data populated on this report is from June 2012-Aug 2012, so the data is already 3 months old and quickly approaching 4 months old.”¹⁰³³ Consequently, Mr. Burtner recommended that CVS needed “to leverage AGI to give us the ability to update the database used for this report [so we] could then decide how often we want to update the database and keep this information more current; we could potentially have the database updated systematically every Saturday night”¹⁰³⁴ It is unclear if the problem was ever addressed.¹⁰³⁵

In 2013, CVS began work on creating a SOM application within its Archer platform.¹⁰³⁶ Archer is an integrated software platform for managing risks and controls on an enterprise-wide basis.¹⁰³⁷ The purpose of the Archer application was “to capture all the due diligence conducted on orders of interest” including data generated using the Store Metrics system.¹⁰³⁸ However, the AGI “stale” data problem persisted into July 2013 as Kelly Baker, who reported to Mr. Burtner,¹⁰³⁹ highlighted when he wrote in an email that “[t]he data snapshot is a 3 month window that is a year old [and any] analysis I make from that data is, for the most part, **irrelevant and pointless.**”¹⁰⁴⁰ Given that the dispensing data used by the Store Metrics application comes from CVS’ own pharmacies, it is inconceivable why the current, refreshed data could not be included in the Store Metrics platform as Mr. Burtner suggested. However, not doing so rendered the application ineffective as a due diligence tool.

4. Negating the Benefits of MicroStrategy and Store Metrics

The MicroStrategy and Store Metrics reports provided useful data to investigate suspicious orders as the DEA expects. However, CVS implemented processes that effectively undercut the usefulness of the tools. This meant that the CVS SOM process, and by extension the anti-diversion program did not prevent CVS pharmacies from receiving orders being diverted and did not meet the required standards for a good compliance program.

¹⁰³² See A. Burtner Deposition at 311:24-312:14 and 315:7--316:5.

¹⁰³³ See Email from A. Burtner to D. Vanelli and P. Hinkle, Attorney Client Privilege – Store Metrics Report, (Dec. 20, 2012), CVS-MDLT1-000109775.

¹⁰³⁴ *Id.*

¹⁰³⁵ See A. Burtner Deposition at 394:15-18.

¹⁰³⁶ See Craig Schiavo, Year-End Review – 2013 (Finalized), 3, (Mar. 4, 2014), CVS-MDLT1-000120580.

¹⁰³⁷ See RSA, RSA ARCHER PLATFORM, <https://www.rsa.com/en-us/products/integrated-risk-management/archer-platform> (last accessed Mar. 10, 2019).

¹⁰³⁸ See Craig Schiavo, Year-End Review – 2013 (Finalized), 3 (Mar. 4, 2014), CVS-MDLT1-000120580.

¹⁰³⁹ See Kelly James Baker Deposition, 23:11-13, (Jan. 24, 2019).

¹⁰⁴⁰ See Email from K. Baker to C. Schiavo, RE: Archer SOM, (Jul 11, 2013), CVS-MDLT1-000078116 at CVS-MDLT1-000078117.

a. Continuing the Previous IRR Process

CVS' continued use of its past process for handling the IRR reports essentially negated the benefits achieved by these tools. CVS persisted in allowing the daily IRR reviewer to maintain overall control of the investigative process by selecting those orders needing a "deep-dive." However, as discussed previously, the daily IRR reviewer had vested interest in keeping flagged orders to a minimum.

The effect of this continued approach to the daily IRR review is readily apparent in the LP Analyst Studies created by Mr. Burtner. The LP Analyst Time Studies identified IRR flagged orders that were sent on for further investigation beyond the IRR. Below is a chart showing a representative twelve days in 2012, which demonstrates how few orders were sent for investigation across Mr. Burtner's five distribution centers:¹⁰⁴¹

SUMMARY OF AARON BURTNER LP ANALYST TIME STUDIES (JUNE-JULY 2012)¹⁰⁴²

Exhibit	IRR Date	Time to Review IRR	# of Orders Investigated
Burtner Ex. 411	6/14/12	55	0
Burtner Ex. 412	6/15/12	30	0
Burtner Ex. 412	6/17/12	35	1
Burtner Ex. 413	6/14/12	60	3
Burtner Ex. 414	7/3/12	15	0
Burtner Ex. 414	7/4/12	20	0
Burtner Ex. 415	7/6/12	35	0
Burtner Ex. 415	7/8/12	40	0
Burtner Ex. 415	7/11/12	35	1
Burtner Ex. 416	7/17/12	50	0
Burtner Ex. 417	8/21/12	30	0
Burtner Ex. 418	9/6/12	15	2

Gary Millikan, who also helped review IRRs beginning in August 2013,¹⁰⁴³ testified that of the flagged orders that appeared on the IRR, he did due diligence on probably less than 5% of the orders.¹⁰⁴⁴ For the remainder (i.e., 95%), he went no further than reviewing the IRR, which provided no additional insights into whether a suspicious order flagged by the algorithms actually involved diversion and consequently should never be shipped to the customer.

¹⁰⁴¹ See A. Burtner Deposition at 329:18-330:3.

¹⁰⁴² See A. Burtner Deposition at 340-366 and Burtner Exh. 500.

¹⁰⁴³ See Gary Millikan Deposition, 41:5-15, (January 11, 2019).

¹⁰⁴⁴ See G. Millikan Deposition at 213:25-214:20; 223:25-224:4; 232:8-14.

b. Ordering from Outside Vendors

Regardless of what set of algorithms and tools CVS used to run its suspicious order monitoring program, the system output never provided the controlled substances compliance team with a complete picture of a CVS store's ordering pattern for hydrocodone. In addition to placing orders with a CVS-controlled distribution center, CVS stores also had the ability to order hydrocodone products directly from an outside vendor (e.g., McKesson or Cardinal).¹⁰⁴⁵

However, even though CVS maintained pharmacy-specific data on every CVS store, the SOM system had no access to that data and no visibility to the outside vendor orders.¹⁰⁴⁶ The IRR did not consider outside vendor orders when determining if an order was suspicious. This "loophole" created issues for CVS including the fact that "[stores] may order a little from both the OV [Outside Vendor] and the DC to stay under the radar" and in one case CVS had "a store, which had a 68,000 hydrocodone pill loss and was placing phone orders to [an] OV."¹⁰⁴⁷

This "loophole" made it impossible for CVS to fulfill its "Know Your Customer" obligations of which it admits being aware of back in 2008.¹⁰⁴⁸ Furthermore, this "loophole," which existed well before the 2012 discussions referenced here,¹⁰⁴⁹ was still not closed as of July 2013;¹⁰⁵⁰ once more illustrating CVS' reluctance to undertake the necessary system improvements to achieve an effective suspicious order monitoring program.

12.6 Accountability - Consistent Enforcement

12.6.1 CVS did not hold employees accountable for failing to maintain and operate an effective anti-diversion program.

There was no evidence presented to demonstrate that CVS held employees accountable for its failed anti-diversion program. In fact, crucial employees, with responsibility for shaping, maintaining and operating CVS' anti-diversion program (e.g., Frank Devlin, John Mortelliti, and Ron Link) continued in positions of substantial authority with CVS even after the failure of its controlled substances compliance program. By failing to hold individuals in substantial authority accountable for the controlled substances program's established lack of effectiveness, CVS's professed commitment to controlled substances compliance is hollow.

¹⁰⁴⁵ See M. Vernazza Deposition at 46:20 to 47:14.

¹⁰⁴⁶ See J. Mortelliti Deposition at 210:17-212:20; see also Email from P. Hinkle to F. Devlin, Conference Call Notes - 10 5 12.docx, (Oct. 5, 2012) ("All orders generated for Outside Vendors are not pushed through the SOM process."), CVS-MDLT1-000033580.

¹⁰⁴⁷ See Email from C. Schiavo to T. Bourque, Monday Morning Meeting w/ Tom P, (Jan. 18, 2013), CVS-MDLT1-000103329.

¹⁰⁴⁸ See M. Vernazza Deposition at 148:5-14; see also Discussion *infra* (concerning CVS Stores #3322 and #4800).

¹⁰⁴⁹ See M. Vernazza Deposition at 46:20 to 47:14; D. Vanelli Deposition at 75:16-18. While no start date is specified, the testimony of both Messrs. Vernazza and Vanelli seem to indicate that CVS stores always had this ability.

¹⁰⁵⁰ See D. Vanelli, *Logistics Planning Update*, 3 (Jul. 8, 2013) ("SOM process will include store controlled substances orders placed with ... outside vendors."), CVS-MDLT1-000100362.

13 Walgreens Boots Alliance, Inc.

13.1 Background

Walgreens Boots Alliance, Inc. (“Walgreens” or “WBA”) began in 1901 as single pharmacy on Chicago’s southside owned and operated by Charles R. Walgreen, Sr.¹⁰⁵¹ Since 1901, WBA has grown into a global company with sales of \$131.5 billion for FY 2018.¹⁰⁵² Formed in 2014 from the merger of Walgreens and Alliance Boots, WBA combined the largest U.S. drugstore chain with Europe’s largest pharmaceutical wholesaler.¹⁰⁵³ As of the end of FY 2018, Walgreens employed more than 415,000 people, and the combined company operated over 18,500 stores in 11 countries under the banners of Walgreens, Duane Reade, Boots and Alliance Healthcare.¹⁰⁵⁴ In addition, WBA maintains a pharmaceutical wholesale and distribution network that includes over 390 distribution centers.¹⁰⁵⁵ Within the U.S., Walgreens’ employs more than 240,000 people and currently maintains more than 9,500 stores.¹⁰⁵⁶

Walgreens, in March 2013, entered into a series of agreements with AmerisourceBergen that has resulted in WBA owning, by the end of FY 2017, 26% of AmerisourceBergen’s common stock in exchange for a ten-year pharmaceutical distribution agreement.¹⁰⁵⁷ In 2015, WBA attempted to acquire Rite Aid, but by 2017 in the face of Federal Trade Commission (“FTC”) pressure, WBA chose instead to acquire 2,186 (almost 50%) of Rite Aid stores.¹⁰⁵⁸

From a corporate compliance perspective, Walgreens has had a Chief Compliance Officer since 1999 when Chester G. Young was appointed to the role for Walgreen Co.¹⁰⁵⁹ Since that time Walgreens has had three

¹⁰⁵¹ WALGREENS BOOTS ALLIANCE, INC., *History*, <https://www.walgreensbootsalliance.com/about/history/> (last visited Feb. 20, 2019). It should be noted that the defendants in this matter are Walgreens Co. and Walgreens Eastern Co, both of which are subsidiaries of WBA.

¹⁰⁵² See Walgreens Boots Alliance, Inc., *Annual Report 2018*, 1 (Oct. 24, 2018) [“WBA 2018 Annual Report”].

¹⁰⁵³ See Ellen Jean Hirst, *Walgreen-Alliance Boots deal is complete*, Chicago Tribune (Dec. 31, 2014, 9:30 AM), <https://www.chicagotribune.com/business/ct-walgreen-completes-merger-0101-biz-20141231-story.html>.

¹⁰⁵⁴ *Id.*

¹⁰⁵⁵ *Id.*

¹⁰⁵⁶ See WALGREENS, *Facts & FAQs*, <https://news.walgreens.com/fact-sheets/frequently-asked-questions.htm> (last visited Feb. 7, 2019).

¹⁰⁵⁷ See Walgreens Boots Alliance, Inc., *Annual Report 2017*, 1-2 (Oct. 24, 2017).

¹⁰⁵⁸ See David Goldman, *A drug store deal gone bad: Walgreens merger with Rite Aid falls apart*, CNN BUSINESS (Jun. 29, 2017, 10:19 AM ET), <https://money.cnn.com/2017/06/29/news/companies/walgreens-rite-aid/index.html>.

¹⁰⁵⁹ See Jim Frederick, *Walgreens veteran Young to retire: Merten stepping into compliance role*, Drug Store News (Nov. 30, 2009) at <https://www.drugstorenews.com/news/walgreens-veteran-young-retire-merten-stepping-compliance-role/>

changes in its Chief Compliance Officer with Laura Merton (2010 to 2013),¹⁰⁶⁰ Diane Nobles (2013-2016),¹⁰⁶¹ and Matthew D'Ambrosio (2017 to present)¹⁰⁶² all serving in the role.

In 2008, Walgreens entered into a five-year Corporate Integrity Agreement (“CIA”) with the Department of Health and Human Services, Office of Inspector General (“OIG”) to settle a series of cases involving healthcare fraud and false claims involving Medicaid reimbursements and the company’s Therapeutic Interchange Program.¹⁰⁶³ Once more in January 2019, Walgreens entered into another five-year Corporate Integrity Agreement and agreed to pay almost \$270 million to settle false claim allegations of improperly billing healthcare programs.¹⁰⁶⁴

From a controlled substances perspective, the relevant time period for this review is 1998 to 2014 during which Walgreens maintained its own internal distribution network that supplied controlled substances to its retail stores.¹⁰⁶⁵ That internal distribution network, however, did not meet all of Walgreens’ retail needs and Walgreens’ stores also fulfilled some of their controlled substances needs from the Group 1 distributors.

By 2012, Walgreens had thirteen (13) distribution centers (“DCs”) capable of distributing controlled substances, and of those, only three (3) handled Schedule II controlled substances (Jupiter, Florida; Perrysburg, Ohio; and Woodland, California).¹⁰⁶⁶ On September 13, 2012, the DEA issued an Order to Show Cause and Immediate Suspension Order (“OTC and ISO”) to Walgreens asserting that the Jupiter distribution center constituted “an imminent danger to the public health and safety.”¹⁰⁶⁷

In February 2013, the DEA issued similar Subpoenas and a Warrant of Inspection on the Perrysburg Distribution Center in Ohio for issues similar to those which the DEA found with the Jupiter DC in Florida and which ultimately led to the Jupiter DC’s license being suspended and its vault being padlocked.¹⁰⁶⁸ Within weeks of receiving the six subpoenas and the warrant, Walgreens decided to “discontinue distribution of controlled

¹⁰⁶⁰ *Id.*; see also Laura Merton LinkedIn Profile, <https://www.linkedin.com/in/lauramerten/> (last accessed Feb. 12, 2019).

¹⁰⁶¹ See Diane Nobles LinkedIn Profile, <https://www.linkedin.com/in/dbnobles/> (last accessed Feb. 12, 2019) (Prior to joining Walgreens, Ms. Nobles served as Senior Vice President, Compliance & Integrity at CVS Caremark).

¹⁰⁶² See Matthew D'Ambrosio LinkedIn Profile, <https://www.linkedin.com/in/mattdambrosio/> (last accessed Feb. 12, 2019).

¹⁰⁶³ See FDAnews Drug Daily Bulletin, Walgreens Settles Investigations Related to Medicaid Reimbursements, FDANEWS (Jun. 6, 2008), <https://www.fdanews.com/articles/107388-walgreens-settles-investigations-related-to-medicaid-reimbursements>; see also Corporate Integrity Agreement Between the Office of Inspector General of the Department of Health and Human Services and Walgreen Co. (Jun. 2, 2008) [“Walgreens 2008 CIA”].

¹⁰⁶⁴ See Beth Jones Sanborn, *Walgreens settles False Claims suits for \$270 million*, HEALTHCARE FINANCE (Jan. 24, 2019), <https://www.healthcarefinancenews.com/news/walgreens-settles-false-claims-suits-270-million>.

¹⁰⁶⁵ However, there is some evidence suggesting that Walgreens had a SOM process before 1998. See, e.g., Email from D. Coughlin to Marcella Ranick, *et al.*, Re: Suspicious Controlled Drug Orders, (August 3, 2010), WAGMDL00660331.

¹⁰⁶⁶ See Walgreens Boots Alliance, *Walgreen Co. Controlled Substance Anti-Diversion and Compliance Program*, 12 (Jul. 17, 2012), WAGMDL00659802 [“WBA CS Program 2012”].

¹⁰⁶⁷ See Letter from Michele Leonhart to Walgreen Company, *Order to Show Cause and Immediate Suspension of Registration* (Sept. 13, 2012), WAGMDL00387653 at WAGMDL00387654 [“Jupiter Show Cause Order”].

¹⁰⁶⁸ See In the Administrative Matter of Walgreens Corporation, (W.D. Ohio 2013), WAGMDL00493697; U.S. Drug Enforcement Administration Subpoenas to Walgreens Corporation, 17-13-7042 (2013), WAGMDL00493694.

substances from the Perrysburg facility” in order to “eliminate any immediate need for further DEA administrative action” regarding the Perrysburg facility.¹⁰⁶⁹

The Jupiter OTC and ISO and the Perrysburg investigation resulted in Walgreens and the DEA entering into a settlement agreement in 2013 to resolve what the DEA noted as “systemic” issues.¹⁰⁷⁰ As part of that settlement, Walgreens agreed to undertake various prospective compliance measures that impacted all of Walgreens’s controlled substances distribution centers.¹⁰⁷¹ Facing mounting pressures and increased enforcement surrounding the distribution of opioids together with signing a long-term distribution arrangement with ABC, Walgreens ceased self-distributing Schedule II controlled substances in 2013. With the reclassification of hydrocodone combination products (“HCPs”) from Schedule III to Schedule II,¹⁰⁷² Walgreens, by October 2014, ended internal distribution of all controlled substances shifting distribution of controlled substances to its outside partner, AmerisourceBergen.¹⁰⁷³

13.2 Executive Summary

The overall theme to the Walgreens’ controlled substances compliance program is “too little, too late.” Even though Walgreens operated licensed distribution centers to supply its stores with controlled substances from 1998, Walgreens made little attempt to design and operate an effective anti-diversion program that identified and reported suspicious orders to the DEA.

That began to change somewhat in 2008 in response to the DEA action taken against Walgreens’s then-primary vendor, Cardinal Health.¹⁰⁷⁴ At that point, Walgreens embarked on an effort to develop an improved and automated anti-diversion program. However, it took several more years (until 2012), and the threatened loss of at least one of its three Schedule II controlled substances distribution centers for Walgreens to begin to take meaningful steps towards implementing a credible controlled substances program. Meanwhile, by March 2013 and the signing of the ABC distribution agreements, Walgreens had decided to cease distributing controlled substances, preferring instead to outsource it to a strategic partner in which Walgreens held a substantial equity stake.

Some of the key contributing factors to this “too little, too late” approach and the failure of Walgreens to take its corporate anti-diversion obligations seriously include:

¹⁰⁶⁹ See Letter from Alice S. Fischer, *et al.*, to C. Lee Reeves, II, *et al.*, Walgreens Perrysburg, Ohio Distribution Center, (Feb. 20, 2013), WAGMDL00674280.

¹⁰⁷⁰ See Settlement and Memorandum of Agreement between the U.S. Department of Justice, U.S. Drug Enforcement Administration and Walgreen Co. (Jun. 10, 2013), WAGMDL00490963 [Walgreens SMOA].

¹⁰⁷¹ See Walgreens SMOA at Addendum: Prospective Compliance (Jun. 10, 2013), WAGMDL00490963 at WAGMDL00490976.

¹⁰⁷² See 79 Fed. Reg. 49661 (Aug. 22, 2014).

¹⁰⁷³ See E. Bratton Deposition at 141:3-4 (Edward Bratton, a Manager in Walgreens Pharmaceutical Integrity, was Walgreens’ 30(b)(6) witness) [“E. Bratton Deposition”]

¹⁰⁷⁴ See, Email from M. Bleser to J. Berkowitz, *et al.*, RE: Update (Apr. 6, 2012) (“In March 2008, a team consisting of Rx Purchasing, Logistics, Legal, Loss Prevention and Store Order System IT began creating the Suspicious Order Monitoring Process in response to the Cardinal incident in which 3 of their DCs were shut down by the DEA for suspicious drug ordering violations”), WAGMDL00709395.

- **Singular Retail Focus:** As one of the top three retail pharmacy chains in the U.S., Walgreens’ efforts to manage controlled substances compliance focused primarily on ensuring its anti-diversion program did not impinge on the retail stores’ ability to obtain the volume of opioid products that the stores requested. Consequently, Walgreens failed to implement effective anti-diversion measures that would regulate or limit a store’s ability to obtain uncontrolled amounts of opioids. This singular retail focus was couched and cloaked under the mantle of “you [have] got to take care of [the] patients,”¹⁰⁷⁵ which merely allowed Walgreens to protect its retail market and justify its noncompliance with its anti-diversion obligations.
- **Lack of Time, Attention & Resources:** Internally, distribution center compliance with SOM and anti-diversion requirements did not receive the time, attention, and resources it deserved. This occurred despite ample available information demonstrating that Walgreens knew or should have known what its obligations as a controlled substances distributor were.¹⁰⁷⁶ It also occurred because the team charged with controlled substances compliance (e.g., Pharmaceutical Integrity, Rx Purchasing, Logistics, Legal, Loss Prevention, & Store Order System IT) did not appreciate that opioids were not “widgets” and thus required special focus at the distribution level.
- **Over-Reliance on Technology:** Within Walgreens, a widespread belief emerged that the Bancroft algorithm and the CSOM system were the solutions to the Walgreens anti-diversion and suspicious order monitoring obligations. Therefore, Walgreens placed undue faith in and reliance on its technologic prowess for inventory management. Consequently, Walgreens did not appreciate that while well-designed technologic solutions were good at identifying outliers, these solutions were not good at resolving those discrepancies. Thus, Walgreens neglected the human element in the compliance equation as demonstrated in the non-IT portions of its program. This faith and overreliance on a technology-driven approach to suspicious order identification and controlled substances compliance were precisely what Joseph Rannazzisi cautioned against in his December 2007 letter to all DEA registrants.¹⁰⁷⁷

When taken together from 1998 to 2014, Walgreens’ controlled substances compliance program was inadequate, and, in my opinion, did not rise to the foundational level on the compliance maturity and program effectiveness model. This failure is particularly troubling given the fact that Walgreens was subject to a Corporate Integrity Agreement during this period, in addition to its DEA settlement, that should have caused the company to take its compliance obligations more seriously than it did.

13.3 Impact

Walgreens activities in Ohio were the predictable result of the company’s failure to maintain a credible anti-diversion program. For example, a suspicious order report (“SOR”) showing store orders from approximately

¹⁰⁷⁵ See, e.g., N. Polster Deposition, 144:2 (Jan. 23, 2019); see also Email from K. Crawford to S. Hasen, *et al.*, FW: OXYCODONE no longer being ordered via PDQ (Oct. 1, 2012) (“We have to do what’s right for patients also.”), WAGMDL00705321. Mr. Crawford was President of Walgreens Health and Wellness division.

¹⁰⁷⁶ See Jupiter Show Cause Order at ¶¶ 7-8.

¹⁰⁷⁷ See Letter from J. Rannazzisi to All Registrants (Dec. 27, 2007).

February to August 2010 nationwide contains 1,712 pages.¹⁰⁷⁸ Below are just a few examples illustrating how Walgreens' anti-diversion program failures translated into various Walgreens stores obtaining high levels of opioids with little or no investigation or interrogation.

Walgreens Store 3226

In the case of Store 3226 located at 6410 Broadway Avenue in Cleveland, the SOR report shows that this store in March, April, and June 2010 obtained 2,200, 2,100 and 2,800, dosage units respectively of oxycodone per month when its limit was 1,800 dosage units (600 base dosage units x factor of 3).¹⁰⁷⁹ Thus, the store was allowed by Walgreens to exceed its limit even with a buffer of 1,200 dosage units per month factored in.

Store 3226, in August 2013, also reported that even though its allocation was increased in June, it needed to be increased again.¹⁰⁸⁰ Walgreens' Pharmaceutical Integrity increased the allocation again without taking into account that Store 3226 was repeatedly out of compliance in 2010 and even went so far as to suggest that the store in the interim before the new allocation becomes effective could order additional bottles via the PDQ process.¹⁰⁸¹

Walgreens Store 3314

From 2006-2014, Store 3314, located at 5400 Pearl Road, Parma, Ohio, obtained on average 276,900 dosage units of oxycodone and 259,420 dosage units of hydrocodone per year from both Walgreens distribution centers, as well as Cardinal Health, Anda and AmerisourceBergen.¹⁰⁸² While the annual averages are high, those numbers do not show the complete picture. Store 3314 purchases of oxycodone from 2006 to 2010 reveal a precipitous year-on-year increase as outlined below.¹⁰⁸³

Walgreen Store 3314 Oxycodone Purchase by Year

YEAR	DOSAGE UNITS
2006	168,200
2007	199,000
2008	242,100
2009	299,000
2010	361,100

¹⁰⁷⁸ See Suspicious Drug Control Orders Report (Aug. 2, 2010); WAGMDL00183798 at WAGMDL00183799 ["SOR"]; E. Stahmann Deposition at 290:24 to 291:15 (Oct. 16, 2018).

¹⁰⁷⁹ See SOR at WAGMDL00205380.

¹⁰⁸⁰ See Email from P. Daugherty to J. Whited, RE: Percocet (Aug 9, 2013) (referencing email from Pharmacy Manager 03226 to J. Whited on Aug. 9, 2013), WAGMDL00698150.

¹⁰⁸¹ *Id.*

¹⁰⁸² See Opioid Shipments to BW4673554 by Distributor 2006-2014, Exhibit 13 to Deposition of Eric Stahmann.

¹⁰⁸³ *Id.*

Thus, from 2006 to 2010, the store's annual number of dosage units increased by an astounding 214.68%. However, Walgreens continued shipping oxycodone to the store from its own distribution centers and allowed it to continue purchasing from the other outside distributors without interruption and without the proper due diligence documentation demonstrating a legitimate basis for this increase.

Walgreens Store 12444

Store 12444, located at 3415 Clark, Cleveland, Ohio, submitted a CSO Override form to increase its allotment of oxycodone with acetaminophen 5mg/325mg to 2,000 dosage units per week (8,000 per month).¹⁰⁸⁴ The rationale provided for the increase was that the store averages 600 opioid prescriptions per day because it serves an emergency room, hospital, pain management clinic, and hospice care.¹⁰⁸⁵ Despite the "red flag" indications of potential diversion, Mayur Tailor from Pharmaceutical Integrity approved the increase less than 24 hours later stating that although the store had reached its product allotment, the increase was approved "after reviewing the item movement."¹⁰⁸⁶ Thus, the limit was increased simply to accommodate the sales line, and not because an investigation revealed that diversion was unlikely to occur.

13.4 Company Commitment – Compliance Culture, Organization & Resources

13.4.1 Walgreens was so focused on the retail stores that it resisted and ultimately failed to adopt anti-diversion measures that would regulate or limit a retail store's ability to obtain uncontrolled amounts of opioid products.

Walgreens singular focus on its retail stores was cloaked by the mantra of "you have to take care of the patient." As a result of this retail focus culture, it was not until the company was forced to confront the magnitude of its noncompliant situation with the Jupiter Order to Show Cause, that Walgreens began making substantive changes to its controlled substances compliance program:

[T]he Company has enhanced its suspicious order monitoring program for controlled substances in an effort to convince DEA that the proposed penalty is excessive and that our new processes will ensure that similar incidents do not recur.¹⁰⁸⁷

For example, when describing Walgreens' pre-2013 suspicious order system, Natasha Polster, former head of the Pharmaceutical Integrity department stated, "the system was **designed** and built way back in the day to ensure that the store was able to get the product in that they need **to take care of their patients**."¹⁰⁸⁸ Even after

¹⁰⁸⁴ See Email from J. Whited to RxIntegrity, Controlled Substance Order Quantity Override Form, (Jun. 18, 2013 at 3:18 PM), WAGMDL00698296 at WAGMDL00698297.

¹⁰⁸⁵ *Id.*

¹⁰⁸⁶ See M. Tailor to J. Whited, RE: Controlled Substance Order Quantity Override Form, (Jun. 19, 2013 at 9:25 AM), WAGMDL00698296.

¹⁰⁸⁷ See Email from T. Polster to D. Doyle, FW: Proposed org chart, at 2 (Dec. 16, 2012) (Doyle was Vice President of Finance), WAGMDL00659270.

¹⁰⁸⁸ See N. Polster Deposition at 161:11-14 (emphasis added).

the Bancroft algorithm was in use, Ms. Polster stated that the system “was designed so we could ensure that the number of tablets ... that goes into any store makes sense for the peer group and business of that store.”¹⁰⁸⁹

The Pharmaceutical Integrity Department, which as of 2012 was charged with overseeing Walgreens’ SOM system, viewed the SOM system as an inventory control mechanism rather than as a compliance control mechanism:

Q: Now, Walgreens' system, similar to my alarm, is there to detect a potential red flag. Would you agree with that?

A: It was put in place to **ensure that the stores had the proper quantities. Not necessarily to ... detect a red flag.** The whole idea was to make sure that the stores were getting the quantities that they needed based on their peer group.¹⁰⁹⁰

Consequently, from an inventory management perspective, “the whole point behind it [the system] was to have simplicity,”¹⁰⁹¹ instead of focusing on Walgreens’ anti-diversion obligations. Denman Murray, Director of Rx Supply Chain Retail, echoed this when he said in his deposition, “traditionally, we’ve always treated a controlled substance like any other, **[a] widget’s a widget to the system.**”¹⁰⁹²

13.4.2 From 2008 to 2013, Walgreens has failed to integrate its controlled substance compliance efforts with the corporate compliance program in any meaningful way.

With its 2008 Corporate Integrity Agreement, Walgreens agreed to continue its voluntary compliance program, including maintaining both a Chief Compliance Officer and a Compliance Committee for a period of five-years (through at least June 2013).¹⁰⁹³ Walgreens also agreed to maintain its Pharmacy Code of Conduct.¹⁰⁹⁴

A. Codes of Conduct

Walgreens currently maintains two separate and distinct Codes of Conduct. The first is the WBA Code of Conduct and Business Ethics (“Business Ethics Code”), and the second is the Walgreens’ Pharmacy and Healthcare Professionals Commitment to Compliance (“Pharmacy Code”).¹⁰⁹⁵ The Pharmacy Code, formerly

¹⁰⁸⁹ See *id.* at 175:4-7. For a description of the Bancroft algorithm, see section 8.3.2 below.

¹⁰⁹⁰ See *id.* at 223:13-23 (emphasis added).

¹⁰⁹¹ *Id.* at 175:10-11.

¹⁰⁹² See D. Murray Deposition, 31:20-22 (Jan. 15, 2019) (emphasis added).

¹⁰⁹³ See Walgreens 2008 CIA at §§ I and III.A.

¹⁰⁹⁴ See *id.* at § III.B (The full title of the document was “Walgreens Pharmacy and Health Care Code of Conduct Policy”).

¹⁰⁹⁵ See Walgreens Boots Alliance, *Code of Conduct & Business Ethics* (Nov. 2017), <https://investor.walgreensbootsalliance.com/static-files/b618a6e0-100d-4f35-a980-57c2c36089b1> [“Business Ethics Code”]; Walgreens, *Pharmacy and Healthcare Professionals Commitment to Compliance*, Policy WAG-POL-PHA-001, Version 1.0 (Feb. 16, 2017), WAGMDL00254921[“Pharmacy Code 2017”].

known as the “Walgreens Pharmacy and Health Care Code of Conduct and General Training,” traces its origins back to July 2008 and is the Code referenced in the 2008 Corporate Integrity Agreement.¹⁰⁹⁶

Despite having two codes of conduct, Walgreens has not expressly linked them together. For example, while the Business Ethics Code is the responsibility of Walgreens’ Chief Compliance Officer, the Pharmacy Code is “owned” by the Senior Vice President of Pharmacy and Retail Operations. The Codes also have two different foci.

The Business Ethics Code “defines how you should conduct yourself as an employee or representative of WBA” and it addresses the responsibilities of WBA officers and employees “to each other, and to customers, suppliers, consumers, and governments.”¹⁰⁹⁷ WBA positions the Business Ethics Code as “a resource to be used to help guide your actions and provides details on **where to go for more information on a particular subject**, to ask questions, or to report a problem.”¹⁰⁹⁸

Under the Business Ethics Code, compliance with controlled substances requirements is not specifically highlighted. The closest the Business Ethics Codes comes to addressing that “particular subject” is located in the “A Foundation of Trust for Our Communities—We comply with healthcare laws” section that was expanded in October 2015.¹⁰⁹⁹ The Business Ethics Code states:

WBA is committed to full healthcare law compliance internationally. All businesses must comply with all laws relating to the commercialization and distribution of healthcare products and the conduct of business in the healthcare industry.¹¹⁰⁰

The section contains no reference to the Pharmacy Code or to the Senior Vice President of Pharmacy and Retail Operations. Instead if employees have questions, they are directed to contact their “manager, legal department or the Global Chief Compliance and Ethics Officer.”¹¹⁰¹

The Pharmacy Code, on the other hand, “covers the expected conduct and ethical principles for all Walgreens Family of Companies team members who work in the pharmacy, Healthcare Clinic or handle any prescription drugs.”¹¹⁰² It specifically includes distribution center personnel and mandates compliance with requirements

¹⁰⁹⁶ See, generally Pharmacy Code 2017; see also Walgreens 2008 CIA at § III.B. The Pharmacy Code has been revised multiple times since 2008 (e.g., 2009, 2010, 2011, 2012, and 2013). A more detailed discussion of those pre-2017 versions can be found in later in the report). Discussion and reference to the Pharmacy Code in this section is limited to the 2017 version (Policy WAG-POL-PHA-001, Version 1.0).

¹⁰⁹⁷ See Business Ethics Code at 6.

¹⁰⁹⁸ *Id.* (emphasis added).

¹⁰⁹⁹ See Business Ethics Code at 31; see also Walgreens Boots Alliance, Notice Regarding Update Walgreens Boots Alliance, Inc. Code of Conduct and Business Ethics (Oct. 20, 2015), <https://investor.walgreensbootsalliance.com/static-files/90962a02-97b7-48b7-b849-4e26c5ed0b96>.

¹¹⁰⁰ See Business Ethics Code at 31. Prior to the October 2015 expansion, the 2014 Walgreens Code of Business Conduct had a much narrower focus addressing only “clinical and regulatory standards” and laws “designed to prevent, detect and punish fraud, waste and abuse.” See Walgreens, Walgreens Code of Business Conduct, 19 (Oct. 2014) at <https://www.readkong.com/page/walgreens-code-of-business-conduct-8174589>.

¹¹⁰¹ *Id.*

¹¹⁰² See Pharmacy Code 2017 at 1.

“Preventing and Mitigating Diversion of Controlled Substances.”¹¹⁰³ The Pharmacy Code, furthermore, does not state expressly that it is a subset of or in addition to the Business Conduct Code. Consequently, Pharmacy and Health Care Professionals could view the Pharmacy Code as the only Code of Conduct they need to follow.

The maintenance of two separated and unlinked Codes of Conduct increases complexity and the likelihood that the two documents will become out of sync. By way of illustration, the Business Code refers to the “Global Chief Compliance and Ethics Officer,” while the Pharmacy Code refers to the “Global Chief Compliance and Privacy Officer.”¹¹⁰⁴ Although they both date to 2017, they are already out of sync with one another. It simply is not leading practice and should not happen especially when the documents are part of a crucial government commitment, such as a Corporate Integrity Agreement.

B. Organization

Like the separate Codes of Conduct, there is no apparent linkage between those responsible for controlled substances compliance and the Chief Compliance Officer’s team. Walgreens 2008 CIA specified that the Compliance Officer “shall be a member of senior management of Walgreens, shall make periodic (at least quarterly) reports regarding Federal health care program compliance matters directly to the Audit Committee of the Board of Directors of Walgreens, and shall be authorized to report on such matters to the Audit Committee of the Board of Directors at any time,” but “not be or be subordinate to the General Counsel or Chief Financial Officer.”¹¹⁰⁵ However, a 2012 chart showing Walgreens Corporate Organization does not show the Chief Compliance Officer as a direct report of Walgreens President and CEO, Greg Wasson.¹¹⁰⁶ In fact, the Corporate Compliance function is not specifically listed on the chart.

On the same Corporate Organization chart, Compliance and Loss Prevention are shown under the Human Resources Division.¹¹⁰⁷ From this chart, it is unclear whether “compliance” encompasses the Chief Compliance Officer’s function but placing Compliance and Loss Prevention under the Human Resources professional staff function is both unusual and I believe a clear indicator that Walgreens did not value these functions sufficiently.

Likewise, Walgreens placed the Pharmaceutical Integrity Department many layers down in the organization under the Pharmacy Operations branch of the Health and Wellness business unit is a clear indication that the department is viewed by Walgreens senior management as relatively unimportant.¹¹⁰⁸ Situated where it was in a business unit, Pharmaceutical Integrity had no meaningful organizational connection either with the Corporate Compliance department or the Corporate legal team, both of whom reside under the professional staff functions. Therefore, it follows the prior discussion that Walgreens viewed controlled substances compliance as

¹¹⁰³ *Id.* at 2-3.

¹¹⁰⁴ See Business Ethics Code at 31; Pharmacy Code 2017 at 6.

¹¹⁰⁵ See Walgreens 2008 CIA at § III.A.

¹¹⁰⁶ See Org. Chart, Walgreens Corporate Organization (Jun. 5, 2012); WAGMDL00387629.

¹¹⁰⁷ *Id.*

¹¹⁰⁸ See Rex Swords CV (undated), P-WAG-02115 (Swords Deposition Exhibit 1).

an inventory management function (pharmacy operations) and not a true compliance function: “[t]hey would review and monitor orders and dispensing habits of the pharmacy ... [s]o they were supporting the stores.”¹¹⁰⁹

13.4.3 Walgreen’s failures to designate a “high-level” individual or group with sole responsibility for controlled substances compliance or provide enough resources for the group contributed to its ineffective and dysfunctional anti-diversion program.

A. Prior to September 2012

Prior to 2012, responsibility for controlled substances compliance by the distribution centers was not vested in a single group or department, but it was spread across multiple departments as Walgreens represented in an organization chart from mid-2012.¹¹¹⁰ At that time, the SOM team was made up of members from Rx Purchasing, Logistics, Legal, Loss Prevention, & Store Order System IT.¹¹¹¹

Not only was responsibility divided among a group, but the group leadership or “ownership” for controlled substances compliance was in dispute. In April 2012, Ed Svihra, Director of Healthcare Loss Prevention wrote:

To clarify, **Loss Prevention has not been responsible for reporting and taking action with district supervision since January 2011** as you state in your email. We did agree to assist with the analysis and design of the reporting. The Rx Purchasing team still has the responsibility to ensure the proper balance between in-stock condition, while limiting excess on-hand quantities. When the [CS-SOMS] pilot is initiated next month, the role of Loss Prevention will be to address those individuals at store level that may be consistently attempting to manipulate the system to increase orders of controlled substances and pseudoephedrine.¹¹¹²

To which, Denman Murray, Director, Rx Inventory Management Drug Stores, responded, “[t]he Suspicious Order program prohibits ordering of controlled substances outside of tolerance limits. **LP [Loss Prevention] is responsible for monitoring the Suspicious Ordering dashboard.**”¹¹¹³ Likewise Barbara Martin, Manager of Pharmacy Inventory Control under Mr. Murray, maintained that although she was involved with the SOM team and the program beginning in late 2008, she only provided general input on the inventory flow and data in flagged reports, but was not responsible for interpreting the data or taking any action on them.¹¹¹⁴

According to Ms. Martin, the ownership issue ultimately came down to budgets:

¹¹⁰⁹ See Rex Swords Deposition, 62:18-22 (Dec. 21, 2018).

¹¹¹⁰ See Walgreens Presentation, *Walgreen Co. Controlled Substance Anti-Diversion and Compliance Program*, (July 17, 2012) (from speaker’s notes) WAGMDL00659802 at WAGMDL00659818.

¹¹¹¹ *Id*

¹¹¹² See Email from E. Svihra to M. Bleser, *et al.*, Update – Response From Healthcare LP (Apr. 9, 2012), (emphasis added) WAGMDL00580316.

¹¹¹³ See Email from D. Murray to M. Bleser and B. Martin, RE: Update-Response From Healthcare LP (Apr. 9, 2012) (emphasis added), WAGMDL00580316.

¹¹¹⁴ See B. Martin Deposition at 23:20 to 24:3, 74:16-20, 78:1-18 (Jan. 25, 2019).

when you have a number of different teams working on a project like this where we have LP and legal and our group, it just comes down who wants to be the ones to put the money in their budget to have to pay for it.¹¹¹⁵

Therefore, by defusing responsibility and accountability to an informal working committee, Walgreens abrogated its responsibilities to maintain and operate an effective anti-diversion program thus demonstrating a lack of commitment to controlled substances compliance, as well as poor corporate governance.

B. Formation of the Pharmaceutical Integrity Department

It was not until late 2012 with the formation of the Pharmaceutical Integrity Department under the direction of Rex Swords and Natasha Polster that Walgreens started vesting responsibility for controlled substances in a single group.¹¹¹⁶ The Department was created in response to the September 2012 Jupiter OTC and ISO in order “to convince DEA that the proposed penalty is excessive.”¹¹¹⁷ However, despite the seriousness or the urgency of the situation, Walgreens failed to provide that organization with the authority and resources to effectively operate its anti-diversion program.

As previously discussed, the Pharmaceutical Integrity Department was effectively buried in the pharmacy operations function within Walgreens.¹¹¹⁸ Thus it lacked the necessary authority to administer the SOM program or to effect change.

Consistent with this lack of authority, Walgreens failed to resource the function appropriately to meet the demands of operating and maintaining the SOM program. Although a December 2012 pilot of the new automated SOM system revealed that there might be more than 14,000 “flagged” or suspicious orders generated per week,¹¹¹⁹ the Pharmaceutical Integrity Department numbered less than 5 FTEs.¹¹²⁰ At its height, it numbered 11 FTEs.¹¹²¹ Taking into account the anticipated workload of “flagged” orders alone, Walgreens severely under-resourced the department setting it up for failure. Two years later Walgreens was no longer self-distributing controlled substances, and the SOM part of Pharmaceutical Integrity’s distribution center SOM mission was moot.

¹¹¹⁵ *Id.* at 154:11-15.

¹¹¹⁶ *See* N. Polster Deposition at 14:21 to 15:17.

¹¹¹⁷ *See* Email from T. Polster to D. Doyle, FW: Proposed org chart, at 2 (Dec. 16, 2012) (Doyle was Vice President of Finance), WAGMDL00659270.

¹¹¹⁸ *See* above at section 8.3.1(B)(2).

¹¹¹⁹ *See* Email from T. Polster to D. Doyle, FW: Proposed org chart, at 2 (Dec. 16, 2012)

¹¹²⁰ *See* N. Polster Deposition at 240:3-8.

¹¹²¹ *Id.* at 240:9-15.

13.5 Program Core – Requirements, Education, Detection & Corrections

13.5.1 From 1998 to 2014, Walgreens operated an anti-diversion program that lacked the fundamental controls of even a foundational compliance program.

From 1998 to 2009, Walgreens represented that the company had an effective anti-diversion program, however, Walgreens cannot demonstrate that it had even rudimentary suspicious order controls in place during that timeframe. This was highlighted by the Walgreens Internal Audit function in 2008 during an audit of DEA Compliance at the Perrysburg Distribution Center.¹¹²² The auditors uncovered system-wide deficiencies involving “suspicious controlled drug order processing and reporting,” “controlled drug reporting specifically receiving record information,” and a “lack of formalized CII controlled substance policies and procedures.”¹¹²³

Beginning in 2009, “the Company ... enhanced its suspicious order monitoring program for controlled substances in an effort to convince DEA that the proposed penalty is excessive and that our new processes will ensure that similar incidents do not recur.”¹¹²⁴ While it is true that by 2009 Walgreens began devoting more effort, including some additional resources, towards designing, operating and maintaining its SOM process and anti-diversion program, that effort failed to create either an effective or even credible program for controlled substances compliance. This was a systemic breakdown, and Walgreens in virtually every program core area examined was unable to produce verifiable evidence to support its claim that it was operating a credible anti-diversion program.

A. Written Standards

1. Handling Suspicious Drug Orders

The Handling Suspicious Drug Order “policy,” which dates to 1998, was the earliest applicable document produced describing controlled substances compliance requirements.¹¹²⁵ From 1998 to 2012, this appears to be the main governance document for Walgreens’ controlled substance program.¹¹²⁶ Walgreens also admitted it did not have documentation resembling a policy and procedure manual for suspicious orders or diversion prevention.¹¹²⁷ Instead, Walgreens relied on a series of emails, presentations and other informal, uncontrolled documentation such as business requirements documents to detail its compliance efforts.¹¹²⁸

¹¹²² See Memorandum from L. Dettmer, *et al.* to S. Kneller and D. Coughlin, DEA Compliance – Perrysburg Distribution Center (Dec. 22, 2008); WAGMDL00757148 at WAGMDL00757163 [Perrysburg DC Audit].

¹¹²³ *Id.*

¹¹²⁴ See Email from T. Polster to D. Doyle, FW: Proposed org chart, at 2 (Dec. 16, 2012) (Doyle was Vice President of Finance), WAGMDL00659270.

¹¹²⁵ See Walgreens, *Handling Suspicious Orders* (Feb. 15, 2005) (the revised policy notes that it originated on Sept. 8, 1998), WAGFLDEA00001854 and WAGFLDEA00001855.

¹¹²⁶ See E. Bratton 30(b)(6) Deposition at 44:24 to 45:7.

¹¹²⁷ See E. Bratton 30(b)(6) Deposition at 25:20-24.

¹¹²⁸ See *id.* at 53:18-20 (“Three e-mails and one policy section.”).

In July 2012, Walgreens claimed to have several policies and procedures to combat diversion.¹¹²⁹ Walgreens identified those written standards as the:

- Code of Conduct,
- Controlled Substances Prescriptions and Good Faith Dispensing Policy,
- Controlled Substances Pick Up and Inventory Policies and Procedures,
- Customer Authentication Policy, and
- Handling Suspicious Drug Orders Policy.¹¹³⁰

Of those listed standards, only the Code of Conduct, Customer Authentication and Handling Suspicious Drug Orders standards applied to its Distribution Centers.¹¹³¹

This lack of documentation not only was contrary to the requirements for credible controlled substances and corporate compliance programs, it was contrary to industry guidelines as well. HDMA, in its 2008 voluntary industry guidelines, “recommended that, to implement these Industry Compliance Guidelines, **specific written company SOPs** be developed and maintained.”¹¹³² However, other than this single policy governing controlled substances compliance, my review uncovered nothing else resembling a policy or procedure.

The Handling Suspicious Drug Order “policy” was remarkable for its brevity and does not meet even the most basic anti-diversion requirements.¹¹³³ It also failed to provide enough specifics to direct Walgreens staff members on what they were supposed to do.

For example, the “policy” stated that the Logistics and Planning Department sent Suspicious Order Reports to the Distribution Centers.¹¹³⁴ The Distribution Centers were to file the reports for five years and make loss and theft reports to the DEA using DEA Form 106.¹¹³⁵ However, beyond the generic organizational designations, the “policy” did not outline, who specifically within the Logistics and Planning Department or the Distribution Centers was responsible for ensuring Suspicious Order Reports are sent, filed, and any necessary reports made to the DEA. Nor was there any indication of who authored or approved the document.

The “policy” also did not outline the criteria the Logistics and Planning Department used to determine an order was suspicious. Key terms such as “unusual size” and “unusual frequency” were undefined in the document. It did not even incorporate the DEA regulatory requirements by reference to allow a staff member to research the DEA’s definitions of those terms.

¹¹²⁹ See Walgreens Presentation, *Walgreen Co. Controlled Substance Anti-Diversion and Compliance Program*, 5 (July 17, 2012) WAGMDL00659802.

¹¹³⁰ *Id.*

¹¹³¹ *Id.*

¹¹³² See HDMA, *Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances*, 14 (2008) (emphasis added), WAGMDL00673706 at WAGMDL00673708.

¹¹³³ See Walgreens, *Handling Suspicious Orders* (Feb. 15, 2005). The text of the policy is reproduced in Appendix G at Figure 1.

¹¹³⁴ *Id.*

¹¹³⁵ See *id.*

The Distribution Center filing requirements also did not adhere to the basic controlled substances requirements in force at that time. According to the “policy,” the Distribution Centers were only required to notify the DEA about substantial losses or thefts. It did not mandate that the Distribution Center report all suspicious orders to the DEA field office.¹¹³⁶

Also, in contravention to accepted compliance practice at the time¹¹³⁷ and specifically highlighted by the DEA in 2012,¹¹³⁸, there was no requirement that Distribution Centers investigate the circumstances reported to it by the Logistics and Planning Department or take any affirmative action to prevent further losses, thefts, or suspicious orders such as holding current or even future orders to those customers. Therefore, the “policy” failed to mandate what should occur when the Distribution Center encounters “suspicious orders” that did not rise to the level of a substantial loss or a theft.

In April 2012, the Handling Suspicious Drug Orders “policy” was amended to include the following statements:

Effective calendar year 2012, the Controlled Substance Order Monitoring and Prevention System prevents suspicious control drugs from being shipped to the stores. In calendar year 2012, because of the program mentioned, suspicious control drug reports are no longer generated as their shipment is prevented by the system.¹¹³⁹

Like the prior 1998 and 2005 versions, the updated 2012 version of the “policy” did not mandate that the Distribution Center report all suspicious orders to the DEA field office.¹¹⁴⁰ Nor, did it require the Distribution Centers to investigate the circumstances reported to it by the Logistics and Planning Department or to take any affirmative action to prevent further losses, thefts, or suspicious orders such as holding current or even future orders to those customers. All that was required was for the Distribution Centers to notify the DEA about substantial losses or thefts. Therefore, the amended “policy” still did not meet even the most basic anti-diversion requirements of the DEA.

2. Pharmacy Code

Walgreens’ Pharmacy and Health Care Code of Conduct Policy (Pharmacy Code), originally enacted in 2008, applied to all employees working in any pharmacy or pharmacy operations, including those handling-controlled

¹¹³⁶ See 21 C.F.R. § 1301.74(b).

¹¹³⁷ See Letter from Wendy Goggin to John Gray (Oct. 17, 2008) WAGMDL00673706. At the time, Ms. Goggin was DEA Chief Counsel and Mr. Gray was President and CEO of the Healthcare Distribution Management Association. The letter discusses HDMA’s voluntary industry guidelines, “Industry Compliance Guidelines: Reporting Suspicious Orders and Preventing Diversion of Controlled Substances.”

¹¹³⁸ Letter from J. Rannazzisi to All Registrants (Jun. 12, 2012).

¹¹³⁹ See Walgreens, *Handling Suspicious Orders* (Apr. 4, 2012), WAGFLDEA00000027.

¹¹⁴⁰ *Id.*

substances.¹¹⁴¹ Employees were expected to acknowledge their understanding and commitment to the Code annually and it was an element of their performance reviews.¹¹⁴²

The Code, itself, specifically addressed controlled substances providing:

Walgreens is committed to complying at all times with all applicable controlled substances laws and regulations. Pharmacy and other health care team members must read and be familiar with Walgreens Policies and Procedures for a more detailed explanation of such requirements. Distribution center team members must read and be familiar with Walgreens Procedure Manual, CM-15 (Distribution Center Procedures for Handling Controlled Drugs).¹¹⁴³

The Code also stated that “Walgreens is committed to cooperating with law enforcement officials consistent with company policy and with its obligations under applicable state and federal law” and that:

Walgreens WILL NOT TOLERATE an illegal, unprofessional or unethical act by any team member, INCLUDING, BUT NOT LIMITED TO, THE UNAUTHORIZED SALE, POSSESSION, USE OR DIVERSION OF CONTROLLED SUBSTANCES. Team members who divert, or illegally possess, sell, or use controlled substances, or fail to report drug diversion, will subject themselves to state or federal prosecution and disciplinary action up to and including termination, Team members should bear in mind that violating any Controlled Substances Act can lead to arrest, prosecution and/or disciplinary action by the state board of pharmacy, including fines, probation, suspension or revocation of license.¹¹⁴⁴

Employees were obligated to report, in a timely manner, violations of the Code, Walgreens policies and procedures, and diversion including “any suspicious activities involving controlled substances.”¹¹⁴⁵ Although this appeared robust on paper, with Walgreens paucity of policies and procedure, lack of training and no definition of “any suspicious activities involving controlled substances,” it was unlikely to be triggered in the case of controlled substances and thus was ineffective.

3. Customer Authentication Policy

Dating to late 2009, the Customer Authentication Policy outlined the Walgreens pharmacy authentication process, a component of Know Your Customer. The one-page policy had only three main requirements:

- Ensure customers have a valid license from the State Board of Pharmacy and the DEA;

¹¹⁴¹ See Walgreens, Pharmacy and Health Care Code of Conduct Policy, (May 2011) (The origination date of the Pharmacy Code was July 23, 2008), WAGFLDEA00000347.

¹¹⁴² *Id.*

¹¹⁴³ See Walgreens, *Walgreens Pharmacy and Health Care Code of Conduct and General Training*, 1 (May 6, 2011), WAGFLDEA00000127; see also Walgreens, *Walgreens Pharmacy and Health Care Code of Conduct and General Training*, 1 (June 6, 2012), WAGMDL00444056. Since the document contains no revision history indicating what changed, it appears that Walgreens made only minor changes between the 2011 and 2012 versions.

¹¹⁴⁴ *Id.* at 1-2 (emphasis in the original).

¹¹⁴⁵ *Id.* at 2.

- Ensure subsequent orders for controlled substances are “regularly and systematically reviewed ... [and] that such orders are consistent with and reasonable for that particular pharmacy's book of business;” and
- Make sure that “[o]rders which may fall outside the usual and customary scope for that particular pharmacy are automatically reduced, identified via an exception report and subject to additional review.”¹¹⁴⁶

Like the Handling Suspicious Orders policy, this policy lacked specifics on how the process works, as well as who was subject to it and had responsibilities under it. In short, as a policy or a procedure, it was deficient and failed to meet even basic compliance standards and expectations.

4. Pharmaceutical Integrity Department SOPs

There was some evidence that the Pharmaceutical Integrity Department attempted to create some written standards resembling SOPs. For example, there was a nine-page document entitled “Suspicious Order Monitoring Program Policy and Procedures for the Pharmaceutical Integrity Team.”¹¹⁴⁷ However, it was unclear when this document was effective and who approved it.

While it did describe in a detailed manner how to pull reports from the various automated systems and use the central monitoring and control dashboard, it was vague on the criteria for how that information was reviewed or how further investigations were undertaken, leaving it to staff members to use “their professional judgment and pharmacy operations experience,” but stated “unusual trends in historical data” or “patterns in drug families in certain markets or communities” should be escalated.¹¹⁴⁸ However, none of these vague parameters were defined in the document. Therefore, it was deficient on its face as a SOM policy or procedure.

B. Detection & Corrections

1. Order Monitoring Controls

From 1998 to 2009, Walgreens claimed Distribution Center personnel were monitoring suspicious orders and that those personnel used a variety of tools. However, the evidence supporting both the active monitoring of orders and the use of specific tools is at best sketchy.

At the outset, it is unclear exactly what role Walgreens expected its distribution centers to fulfill under the SOM and anti-diversion programs. On the one hand, Edward Bratton noted that distribution centers “are an important part of our system to detect and monitor controlled substances ordering.”¹¹⁴⁹ On the other hand, a 2012 Walgreens presentation describing Walgreens’s program stated:

Distribution Centers are not **designed to be a backstop to the pharmacists** who are the front lines; rather, Distribution Centers are **more akin to supply warehouses**. The stores, on the one

¹¹⁴⁶ See Walgreens, Policy & Procedure Customer Authentication, § 4 b,d,e (Apr. 2, 2012), WAGFLDEA00001746.

¹¹⁴⁷ See Walgreens, *Suspicious Order Monitoring Program Policy and Procedures for the Pharmaceutical Integrity Team*, (undated); WAGMDL00395923 at WAGMDL00395928.

¹¹⁴⁸ *Id.* at 9.

¹¹⁴⁹ See E. Bratton Deposition at 119:8-10.

hand, and corporate headquarters, on the other hand, are best equipped to ensure compliance and assist in combatting controlled substance abuse.¹¹⁵⁰

Walgreens described three different tools utilized from 1998 to 2009:

- **RX Questionable Order Qty** – This process was performed by the distribution center personnel. The procedure called for distribution personnel to review orders and contact the pharmacy if there was a “questionable quantity”.¹¹⁵¹ There was no criteria provided about the levels, no outline of what investigations occurred, and no copies of the reports were provided during discovery.¹¹⁵²
- **“Pickers”** – This control involved relying the employees in the controlled substances vault to identify potentially suspicious orders.¹¹⁵³
- **Chemical Handler’s Reports** – These reports “flagged” potentially suspicious orders after applying the “3x” multiplier from the Chemical Handlers Manual.¹¹⁵⁴ There was no discussion of holding and investigating “flagged.” The reports simply identified orders over the limit, which were collected and simply sent to the DEA. These reports were stopped when the CSOM system was deployed.¹¹⁵⁵

Of the three tools, the Chemical Handler’s Reports generated many suspicious orders. For Ohio alone, there were 5,500 suspicious orders from 2006 to November 2012.¹¹⁵⁶ None of these shipments were properly investigated or stopped.¹¹⁵⁷

Neither the excessive quantity query nor the Chemical Handler’s Report froze “flagged” orders or prevented them from ultimately shipping.¹¹⁵⁸ Even when suspicious orders were identified, I saw nothing documented that suggested that Walgreens did more than a cursory further inquiry before the order shipped.¹¹⁵⁹ Walgreens also admitted that the company is unaware of any due diligence occurring on orders flagged by the Chemical Handler’s Report.¹¹⁶⁰

¹¹⁵⁰ See Walgreens Presentation, *Walgreen Co. Controlled Substance Anti-Diversion and Compliance Program*, (July 17, 2012) WAGMDL00659802 at WAGMDL00659817 (emphasis added) (see speaker’s notes).

¹¹⁵¹ See E. Bratton 30(b)(6) Deposition at 83:11-15; *see also*, Walgreens, Rx Questionable Order Qty Procedure, (Dec. 11, 2006), WAGMDL00757788. The procedure was modified in 2010 and again in 2013, when among other things, the name was changed to “Authentication of Prescription Order Policy.” *See* WAGMDL00751822; WAGMDL00749381.

¹¹⁵² *Id.* at 85:11-23 and 95:11-14.

¹¹⁵³ *Id.* at 113:20 to 114:5; *see also*, Walgreens, Rx Questionable Order Qty Procedure, (Dec. 11, 2006), WAGMDL00757788. The procedure was modified in 2010 and again in 2013, when among other things, the name was changed to “Authentication of Prescription Order Policy.” *See* WAGMDL00751822; WAGMDL00749381.

¹¹⁵⁴ *See* E. Bratton 30(b)(6) Deposition at 144-22-24; *see also* Discussion *infra*.

¹¹⁵⁵ *See* E. Bratton 30(b)(6) Deposition at 150:2-5.

¹¹⁵⁶ *Id.* at 164:19-22.

¹¹⁵⁷ *See* Errata to Ed Bratton 30(b)(6) Deposition, Erratum No. 3.

¹¹⁵⁸ *See* E. Bratton Deposition at 65:20 to 66:11 and 181:18 to 182:3.

¹¹⁵⁹ *Id.* at 95:11-14 and 159:20-24; *see also* Jupiter Show Cause Order at ¶¶11-12.

¹¹⁶⁰ *See* Errata to Ed Bratton 30(b)(6) Deposition, Erratum No. 3.

2. Automated Controlled Substance Reporting (CSOM System)

Starting in 2009, Walgreens began work to develop an Automated Controlled Substances Reporting (“CSR”) system “to identify and reduce excessive orders.”¹¹⁶¹ Work on the new system was divided into five phases, with Phase 5 being deployed in November 2012, more than three years after the start of Phase 1.¹¹⁶² During this “gap” before the new system deployed, Walgreens was not meeting its anti-diversion requirements as the DEA outlined in great detail in the Jupiter OTC and ISO.¹¹⁶³

The new system, later renamed the Controlled Substance Order Monitoring system or CSOM, was conceived to be the primary mechanism to monitor, detect, and report suspicious orders. While it was “a bolt-on” system to the strategic inventory management system (“SIMS”),¹¹⁶⁴ it was considered the centerpiece of the Walgreens distributor anti-diversion program. However, as designed and implemented, the CSOM system failed to achieve that objective.

3. The Bancroft Algorithm

The heart of CSOM system was an algorithm developed by Wayne Bancroft and Tracy Morris in 2008.¹¹⁶⁵ As originally conceived in 2008, the Bancroft algorithm was “a methodology for identifying suspicious orders in terms of order size and order frequency.”¹¹⁶⁶ Order size and order frequency were two of the three DEA listed criteria for determining if a controlled substances order was “suspicious.”¹¹⁶⁷ The regulation also made clear that the three DEA criteria are not the only ones that could render an order as “suspicious.”¹¹⁶⁸

The CSOM system deployed in November 2012 used the algorithm from two dimensions – tolerance and ceiling.¹¹⁶⁹ The tolerance was the number of bottles ordered above the store’s average historical orders based on the last 26 weeks of ordering history.¹¹⁷⁰ The ceiling was based on stores with similar volume when compared to the previous six (6) weeks.¹¹⁷¹ According to the algorithm proposal, “[i]f an order is placed on the

¹¹⁶¹ See Walgreens Presentation, *Controlled Substance Ordering – Evolution of Controlled Substances Ordering Process*, 3 (Oct. 11, 2012) (the system later came to be known as the Controlled Substance Order Monitoring and Prevention System), WAGMDL00667936 at WAGMDL00667938. [“Evolution Presentation”]; see also Appendix G at Figure 2.

¹¹⁶² *Id.*

¹¹⁶³ See generally Jupiter Show Cause Order.

¹¹⁶⁴ See D. Murray Deposition at 35:20-21.

¹¹⁶⁵ See Memorandum from W. Bancroft and T. Morris to S. Bamberg, *et al.* DEA Suspicious Order Reporting, 1 (Jun. 23, 2008) (Since Bancroft was the Lead Business Systems Analyst, IT Integration and the mathematical lead, the algorithm has become associated with his name), WAGMDL00624503 [“Bancroft Algorithm Proposal”].

¹¹⁶⁶ *Id.* at 1.

¹¹⁶⁷ See 21 C.F.R. § 1301.74(b). The third enumerated criterion was a substantial deviation from a normal ordering pattern.

¹¹⁶⁸ *Id.* The regulation specifically states that suspicious orders “include” these three criteria clearly indicating that these listed criteria were not a fixed set.

¹¹⁶⁹ See Evolution Presentation at WAGMDL00667940.

¹¹⁷⁰ *Id.* at WAGMDL00667943.

¹¹⁷¹ *Id.* at WAGMDL00667943

DC [Distribution Center] that exceeds its tolerance limit the order is flagged as suspicious.”¹¹⁷² The final system was being developed to simply reduce orders to whichever dimension yields the lowest value.¹¹⁷³

Although first deployed in August 2009 as a pilot program, the CSOM system underwent many changes through 2014.¹¹⁷⁴ These changes overall did not serve to enhance the system, but rather to weaken it.

Despite the apparently advanced algorithm employed by the CSOM system, Walgreens anti-diversion program was plagued by “loopholes” that effectively negated the value of the algorithm and the system.¹¹⁷⁵ Furthermore, these “loopholes” allowed Walgreens to avoid its obligations, in some cases intentionally and systematically, to report suspicious orders and prevent diversion. Therefore, overall the CSOM system essentially became useless as an anti-diversion control.

4. CSOM “Loopholes”

a. Order Intercept (a.k.a. “Cutting” Orders)

One major loophole in the system involved order intercepts or ordering “cutting.” Rakesh Khanna from Store Replenishment and Forecasting described how the order intercept process worked in a memorandum she wrote and circulated in October 2011.¹¹⁷⁶ While explaining the business reason for addressing suspicious orders, Ms. Khanna wrote:

The purpose of this project [to intercept orders] is to create a process to systematically identify and prevent suspicious orders based on a formula used to determine inconsistent (suspicious) ordering patterns for controlled drugs. Any Control Drug Orders that are deemed suspicious will be flagged as suspicious and populated in a file to be sent up centrally to Loss Prevention and Rx services for review/analysis. The order that is flagged as suspicious on the store side will be intercepted and the order qty will be reduced to a non-suspicious (order limits) level. **The item will be reduced to a non-suspicious level in order to prevent suspicious orders from being sent over to the DC. This method will help to insure [sic.] that the DC does not receive suspicious orders from stores and limit the possibility of fines that may be imposed by the DEA.**¹¹⁷⁷

Thus, although the CSOM system was designed to flag store orders that exceeded the tolerance or ceiling limit for a particular scheduled product at the distribution level, the plan now was to intercept orders that fit the definition of “suspicious,” and thus need further investigation and potential reporting to the DEA, and render

¹¹⁷² See Bancroft Algorithm Proposal at 1.

¹¹⁷³ See Evolution Presentation at WAGMDL00667943.

¹¹⁷⁴ See Appendix G at Figure 2.

¹¹⁷⁵ See N. Polster Deposition at 157:9-18.

¹¹⁷⁶ See Email from R. Khanna to K. Provost, *et al.*, DEA Business Reason (Oct. 27, 2011); WAGMDL00119542.

¹¹⁷⁷ See Email attachment *DEA Intercept Suspicious Order* from R. Khanna to K. Provost, *et al.* at 1 (Oct. 27, 2011) (emphasis added); WAGMDL00119542 at WAGMDL00119543; *see also* Memorandum from T. Polster to R. Swords, *Status* (Nov. 30, 2012) (Under the SOM Meetings section it reads “[w]ork group has been put together to begin the determination between a suspicious order and an order of interest.”), WAGMDL00574824 at WAGMDL00574825.

them “non-suspicious.” By altering (“cutting”) the orders during transmission from store to distribution center, the distribution center would receive an order that was not “flagged” and thus, Walgreens’s position was, the DC would not be required to take any action beyond filling the order. This would reduce the number of orders needing investigation from hundreds per week to a more manageable level.¹¹⁷⁸ However, by taking this approach, Walgreens intentionally and systematically avoided its obligations to investigate and report suspicious orders and prevent diversion.

b. “Flagged” Orders Are Not “Suspicious Orders”

A second major loophole involved the Pharmaceutical Integrity department’s interpretation that orders exceeding the ceiling limit and flagged by the system using the Bancroft algorithm were not suspicious. Orders that exceeded the tolerance limit were “flagged [by the system] as suspicious,”¹¹⁷⁹ and under the controlled substances regulations, suspicious orders are to be investigated and reported “when discovered.”¹¹⁸⁰

The Walgreens Pharmaceutical Integrity Department as of 2012, took the position that “flagged” orders were not “suspicious” as defined by DEA regulations because:

Whether it was suspicious or not did not even come into play because we had an algorithm and we told them that they weren't supposed to go over it and if they needed to go over it, they had to supply the necessary information so that we could have the documentation we needed.¹¹⁸¹

As Ms. Polster further elaborated, an order “flagged” by the system was not considered “suspicious” because she “never felt like any pharmacy manager or pharmacist was doing anything nefarious to get more product into the store.”¹¹⁸² At the outset, although the DEA provides regulatory latitude to include another criterion to classify an order as “suspicious,” the feeling that “no pharmacy manager or pharmacist was doing anything nefarious to get more product into the store,”¹¹⁸³ is neither an objective nor valid gauge, for determining whether an excessive order is suspicious and diversionary.

Ms. Polster also stated that orders “flagged” using the Bancroft algorithm were simply “an order that was over the algorithm that was designed for that specific store and that it did not conform to ... what we thought the store needed.”¹¹⁸⁴ Therefore, since the order did not ship in full, it was not deemed “flagged,”¹¹⁸⁵ not deemed an “order of interest,” and not deemed “suspicious.”¹¹⁸⁶

¹¹⁷⁸ See Email from T. Polster to D. Doyle, FW: Proposed org chart, at 2 (Dec. 16, 2012) (Doyle was Vice President of Finance), WAGMDL00659270. The December 1-8, 2012 test of the CSOM system produced 470 “cut orders.” *Id.*

¹¹⁷⁹ See Bancroft Algorithm Proposal at 1.

¹¹⁸⁰ See 21 C.F.R. § 1301.74(b).

¹¹⁸¹ See N. Polster Deposition. at 174:3-8. See section on Controlled Substances Overrides below for a discussion of the “necessary information.”

¹¹⁸² *Id.* at 175:8-10.

¹¹⁸³ *Id.* at 175:8-10.

¹¹⁸⁴ *Id.* at 175:15-16.

¹¹⁸⁵ *Id.* at 187:5-9.

¹¹⁸⁶ *Id.* at 221:11-21.

Although this was merely a game of semantics, the impact was real. Orders flagged using the Bancroft algorithm were in fact “suspicious,” in accordance with the plain meaning of the word, and therefore needed further investigation and potential reporting under the DEA’s requirements. During the week of December 1-8, 2012, the CSOM system in “tracking mode” generated more than 14,000 potential ceiling limit “orders of interest” and “[e]ach order of interest has to be investigated before more product can be added to the store.”¹¹⁸⁷ So there was a real concern that once fully operational the CSOM system could generate “thousands of ‘orders of interest’ per week.”¹¹⁸⁸ At the time, Walgreens understood there was direct correlation between how much they decreased a ceiling level and the amount of work (number of flagged orders) that reduction generated.¹¹⁸⁹ Walgreens’s contention that because it deemed these “flagged orders” not to be either “of interest” or “suspicious” that these orders were then not “suspicious” for purposes of further investigation or DEA reporting is both specious and non-compliant.

c. Use of Outside Distributors & Visibility to Order Ceilings

As originally implemented, the CSOM system allowed Walgreens stores to order up to their ceiling limits as determined by the Bancroft algorithm. If a Walgreens store hit the store ceiling limit or tried to exceed it, the order was not shipped. This process of simply not shipping an order that exceeded the threshold meant that the store only knew there was a problem when the order failed to arrive.¹¹⁹⁰

From 2009 to 2012, the CSOM system only reviewed store orders placed with the Walgreens distribution centers.¹¹⁹¹ Therefore, if a store’s order of controlled substances failed to arrive, they could place an order with an outside distributor, such as Cardinal Health or AmerisourceBergen, and “so they ordered it through the wholesaler [outside distributor].”¹¹⁹²

Pharmaceutical Integrity did not view this situation as a “loophole” but rather as a situation of poor communication “between the support center and the stores to let them know that they’ve reached the limits we set for them.”¹¹⁹³ and “[t]he wholesaler has their own suspicious order responsibilities.”¹¹⁹⁴

As a result of the poor job “of communicating to the stores why they didn’t get their order,”¹¹⁹⁵ a relatively late adjustment to the system was made to provide stores visibility to their order ceilings.¹¹⁹⁶ Therefore, as Ms. Polster described it:

¹¹⁸⁷ See Email from T. Polster to D. Doyle, FW: Proposed org chart, at 2 (Dec. 16, 2012), WAGMDL00659270.

¹¹⁸⁸ *Id.* at 1.

¹¹⁸⁹ See Email from T. Polster to S. Mills, RE: Ceiling Update, (Jan. 4, 2013), WAGMDL00414048.

¹¹⁹⁰ See N. Polster Deposition at 169:1-7.

¹¹⁹¹ See Walgreens Presentation, *Controlled Substances Talking Points*, 2 (Jun. 12, 2012), WAGMDL00077016 at WAGMDL00077017.

¹¹⁹² See N. Polster Deposition at 250:15-16.

¹¹⁹³ See *id.* at 253:4-7.

¹¹⁹⁴ *Id.* at 250:16-17.

¹¹⁹⁵ See *id.* at 252:11-12.

¹¹⁹⁶ See *id.* at 167:20 to 168:22.

We had to train the stores, leave your orders alone. Let the orders generate by themselves based on the algorithm and if you feel that you need more, fill out the controlled substance override form so that we can look at it, make sure we have proper documentation as to the reason why you need it, and then we will place the order for you.¹¹⁹⁷

Sometime in late 2013 or early 2014, Walgreens implemented a solution that allowed the stores to see how many bottles remained before the ceiling was triggered. With that information, if the stores needed more inventory than what the system allowed, they could order directly from the outside wholesaler rather than fill out a Controlled Substances Override Form.

In the case of Walgreens stores ordering from Cardinal, Cardinal at least by January 2013, was supplying Walgreens with a list of all stores that were within 75% or more of their monthly Cardinal accrual to, as Steve Mills put it, “help us [Walgreens] prevent a SOM from occurring.”¹¹⁹⁸

d. PDQ Orders & “Inter-storing”

Another “loophole” in the CSOM system involved PDQ orders.¹¹⁹⁹ These orders allowed a store to place orders for drug products, including controlled substances, that were needed quickly. Prior to October 2012, Walgreens stores could order oxycodone via the PDQ process once they hit their ceiling limit. PDQ orders were not included in the monthly cumulative limits and therefore, “a store could hit the line limit on their weekly CII whs order and then they could create a PDQ order on a daily basis and far exceed the monthly line limit total we were trying to enforce.”¹²⁰⁰

By October 2012, oxycodone PDQ orders were no longer allowed and “[b]y turning these items off of PDQ, the only way for a store to exceed the line limit is for us to put in a manual order. Hence, the Control Substance Order Override Form.”¹²⁰¹

Although the PDQ avenue was blocked, Kermit Crawford, President of Walgreens Health and Wellness pointed out that a store could simply do an interstore transfer.¹²⁰² Until early 2013, interstore transfers occurred with no visibility by Pharmaceutical Integrity.¹²⁰³ However, according to Ms. Polster, interstoring was controlled via the controlled substances invoicing process that required knowledge by Walgreens leadership.¹²⁰⁴ However,

¹¹⁹⁷ See *id.* at 169:9-16.

¹¹⁹⁸ See Email from N. Rausch (Cardinal Health) to P. Holohan (Cardinal Health), RE BW6664381WALGREEN CO. (Jan. 21, 2013), CAH_MDL2804_00783520; see also Email from S. Mills (Walgreens) to P. Holden (Cardinal Health), RE BW6664381WALGREEN CO. (Jan. 18, 2013) (embedded in N. Rausch thread), CAH_MDL2804_00783520.

¹¹⁹⁹ While no official definition of “PDQ” was found, according to Natasha Polster to her it meant “pretty darn quick.” See N. Polster Deposition at 253:21-24.

¹²⁰⁰ See Email from D. Lovejoy to K. Crawford, *et al.*, FW: OXYCODONE no longer being ordered via PDQ (Oct. 1, 2012), WAGMDL00705321.

¹²⁰¹ *Id.*

¹²⁰² See Email from K. Crawford to D. Lovejoy, *et al.*, RE: OXYCODONE no longer being ordered via PDQ (Oct. 1, 2012), WAGMDL00705321.

¹²⁰³ See N. Polster Deposition at 257:14-16 and 258:1-2.

¹²⁰⁴ *Id.* at 257:21-24.

because interstore transfers avoided the CSOM system, the practice effectively undermined the anti-diversion program by allowing stores to secure amounts of controlled substances well in excess of the established CSOM ceiling amount.

Walgreens store 2865 in Modesto, California is a case in point. During a GFD (Good Faith Dispensing) audit in October 2012 it was noted that:

This store's average movement on hydro/APAP 10/325 is 17,500 tabs a week put them over the corporate limit. This changed their ordering habits with Cardinal, which then led to an SOM with them. I submitted a report which was approved by WAG but denied by Cardinal. **As such this location has had a large increase in interstores. This increase in interstores has led to short supplies at other locations in town.** We need to get Cardinal back on shipping this location additional controls.¹²⁰⁵

e. Removing Stores from the CSOM System

During Phase II of the CSOM project, Rakesh Khanna authored a project request estimate to give “Rx Services ... the ability to remove items from the order limitation process or **remove an entire store from the order limit program for a limited amount of time.**”¹²⁰⁶ An earlier project requirements document suggested this ability was being added “to account for stores that may need to order more of an item for a certain amount of time.”¹²⁰⁷ In other words, to circumvent the SOM system.

This option continued to exist after the formation of Pharmaceutical Integrity allowing them to switch stores to “tracking” only.¹²⁰⁸ However, despite Ms. Polster’s assertion that policies and procedures existed to govern this process and define “a limited amount of time,” no supporting documentation was provided nor had Ms. Polster ever seen those policies and procedures.¹²⁰⁹ Therefore, it is a reasonable assumption that the documentation did not exist, especially given Walgreens overall poor documentation practices.

f. Controlled Substances Overrides

Beginning in 2013, Walgreens implemented the process of Controlled Substances Overrides (“CSO Overrides”). The goal of this process was to ensure there was “adequate review before sending in additional inventory” above the system generated ceiling (Bancroft algorithm).¹²¹⁰ The change resulted because “[t]he previous system would continue to send additional product to the store without limit or review which made

¹²⁰⁵ See Email from M. Federico to D. Murray, GFD Audit for store 286 in Modesto, California, 2 (Oct. 31, 2012) (emphasis added). WAGMDL00113808 at WAGMDL00113810. At the time Michael Federico was District 293 Pharmacy Supervisor and Denman Murray was Director of Rx Inventory Management Drug Stores.

¹²⁰⁶ See R. Khanna, *DEA Suspicious Order Item Limits – Phase II Project P-09002*, 1 (Aug. 26, 2009), WAGMDL00492067.

¹²⁰⁷ See Ora Yelvington, *Intercepted/Suspicious Store Orders Project#: P99999 Requirements Document Version 1.0*, 3 (Feb. 2009) WAGMDL00492626.

¹²⁰⁸ See N. Polster Deposition at 262:4-7.

¹²⁰⁹ *Id.* at 262:13 to 263:14.

¹²¹⁰ See Email from E. Bratton to A. Patel, RE: Controlled Substance Order Quantity Override Form (Aug. 19, 2013), WAGMDL00021425.

possible the runaway growth of dispensing of products like oxycodone, that played a roll in the DEA's investigation of Walgreens."¹²¹¹

However, when Walgreens examined the percentage of CSO Overrides that were approved versus those disapproved, they found that for FY 2014, 95.52% of the overrides submitted were approved which increased slightly to 95.63% in FY 2015.¹²¹² This is consistent with the previously discussed examples of Walgreens Stores 3314 and 12444.¹²¹³ Therefore, as these data show, if a CSO Override form was submitted, it was almost guaranteed to be approved. Such high approval rates are not normal and demonstrate that Pharmaceutical Integrity simply was not conducting a robust interrogation and investigation of the information presented.

5. The Net Effect on the CSOM System

The net effect of the changes implemented to the CSOM system between 2009 and 2013 rendered the system essentially useless as anti-diversion control. In 2016, Walgreens prepared a "State of Rx Integrity" presentation examining data from FY 2014 and FY 2015 (October 2013 to October 2015).¹²¹⁴

From 2014 to 2015, the quantity of oxycodone dispensed across all Walgreens stores increased by 9% and the average quantity per prescription increased by 1.34% for 2014 and 1.96% for 2015.¹²¹⁵ While the total quantity of hydrocodone sales decreased by 4%, the average quantity per prescription increased by 1.66% in 2014 and 5.43% in 2015.¹²¹⁶

Against this backdrop, Walgreens also examined the total population of "flagged" orders for FY 2014 and FY 2015, which appear on average to trend consistently at about 4,000 per month:¹²¹⁷

- **Above the Limit:** Orders exceeding the tolerance limit accounted for 74.16% for FY 2014 and 77.48% for FY 2015 of all "flagged" orders.
- **Below the Limit:** Orders reduced to below the tolerance or ceiling accounted for 25.74% for FY 2014 and 21.64% for FY 2015 of all "flagged" orders.

These data imply that Walgreens stores were routinely submitting orders that were "flagged" by the CSOM system and that once "flagged" they stood less than 1 in 4 chance of being reduced. However, for the other 75% of the orders, it is not clear whether they were shipped or simply were not filled.¹²¹⁸ Notwithstanding Walgreens strained interpretation of what constituted a suspicious order, every "flagged" order either should

¹²¹¹ *Id.*

¹²¹² See E. Bratton Presentation, *State of Rx Integrity*, 22 (May 10, 2016); WAGMDL00010887 at WAGMDL00010888.

¹²¹³ See Discussion *infra*.

¹²¹⁴ See E. Bratton Presentation, *State of Rx Integrity* (May 10, 2016); WAGMDL00010887 at WAGMDL00010888.

¹²¹⁵ *Id.* at 6.

¹²¹⁶ *Id.* at 8. Fewer prescriptions with larger volumes would result in fewer sales.

¹²¹⁷ *Id.* at 17.

¹²¹⁸ See N. Polster Deposition at 332:23-334:3.

have been reported to the DEA as “suspicious” or subjected to further investigation to validate that the “flags” were false positives. By failing to do so, Walgreens’ anti-diversion program was not effective.

C. Audits

While there is some evidence that Walgreens conducted audits of its controlled substances distribution centers, these audits seemed to be *ad hoc* events, rather than part of a normal, programmatic audit cycle. I saw no formal policies or procedures evidencing a programmatic approach to internal auditing of its distribution centers that Walgreens’ Internal Audit Department employed.

When done, these audits highlighted significant flaws with Walgreens controlled substances compliance program.¹²¹⁹ However, although the infrequent internal audit identified systemic shortcomings with Walgreens’ controlled drug reporting process, the company failed to address the matters in a timely fashion. For example, in the case of the internal audit of the Perrysburg distribution center, the audit fieldwork was completed in November 2008,¹²²⁰ but Walgreens’ management only committed to take up the problem in May 2009.¹²²¹

D. Education

There is scant evidence that Walgreens undertook any comprehensive efforts to train its employees, even ones in positions of substantial authority for controlled substances compliance, about the company’s obligations surrounding controlled substances and the DEA’s requirements. For example, Edward Bratton, Natasha Polster and Rex Swords all testified that before coming into their respective roles with responsibility for controlled substances compliance, they had operational but no legal or compliance background.¹²²² They also received no formal training but received on-the-job support from Walgreens’ regulatory and legal staff members.¹²²³

On the topic of general employee training, Mr. Bratton testified that he could not identify any specific employee training programs “about opiates and suspicious order monitoring.”¹²²⁴ Furthermore, no training materials supporting such training programs were located.¹²²⁵

¹²¹⁹ See Internal Audit Report, DEA Compliance – Perrysburg Distribution Center, (Dec. 22, 2008), WAGMDL00757148 at WAGMDL00757163.

¹²²⁰ *Id.* at 2 (Background), WAGMDL00757162.

¹²²¹ *Id.* at Summary of Findings, 2, WAGMDL00757160.

¹²²² See E. Bratton Deposition, 17: 4-10 (Nov. 30, 2018) (discussing his full resume which was not produced) [“Bratton 11/30/18 Deposition”]; N. Polster Deposition at 16:2-10; R. Swords Deposition at 92-8-14.

¹²²³ See Bratton 11/30/18 Deposition at 51:8 to 54:1; N. Polster Deposition at 80:9-20; R. Swords Deposition at 96:1-4.

¹²²⁴ Bratton 11/30/18 Deposition at 138:24 to 139:8; *see also* B. Martin Deposition at 65:1-24.

¹²²⁵ *Id.* at 139:8-11.

13.6 Accountability - Consistent Enforcement

13.6.1 Walgreens failed to enforce the standards outlined in the Pharmacy Code and thus there is no real accountability for the program's lack of effectiveness.

Walgreens' Pharmacy Code mandated full compliance with the requirements pertaining to controlled substances distributors and obligate employees to report breaches of the Code.¹²²⁶ However, there was no evidence presented to demonstrate Walgreens enforced the standards in the Code, especially as it pertained employees in positions of substantial authority for controlled substances compliance.

The crucial employees, with responsibility for shaping, maintaining and operating Walgreens' anti-diversion program (e.g., Natasha Polster, Edward Bratton, and Rex Swords) continued in positions of substantial authority with Walgreens after the failure of its compliance program for controlled substances, and the cessation of internal distribution of Schedule II and III products to the retail locations. By failing to hold these individuals accountable for the controlled substances program's established lack of effectiveness, Walgreens' compliance program and its Pharmacy Code are merely words on paper that profess to hold culpable individuals accountable.

14 Mallinckrodt Pharmaceuticals

14.1 Background

Mallinckrodt Pharmaceuticals ("Mallinckrodt") began with the formation of G. Mallinckrodt and Company in 1867 by the three Mallinckrodt brothers, Gustavo, Otto and Edward to supply local pharmacists because it was the only chemical company west of Philadelphia.¹²²⁷ In 2000, Mallinckrodt was acquired by Tyco International's healthcare division, which became Covidien in 2007.¹²²⁸ Mallinckrodt Pharmaceuticals later split from Covidien in 2013 to become "an independent, \$2 billion public company that develops, manufacturers, markets and distributes specialty pharmaceutical products and diagnostic imaging agents."¹²²⁹

Mallinckrodt's publicly stated mission is "managing complexity, improving lives", which is centered around a set of four values: "patient-centric, integrity, innovative and collaborative."¹²³⁰ With over 3,600 employees

¹²²⁶ See Walgreens, *Walgreens Pharmacy and Health Care Code of Conduct and General Training*, 1-2 (May 6, 2011), WAGFLDEA00000127; see also Walgreens, *Walgreens Pharmacy and Health Care Code of Conduct and General Training*, 1-2 (June 6, 2012), WAGMDL00444056.

¹²²⁷ See MALLINCKRODT PHARMACEUTICALS, *Our Story*, <http://www.mallinckrodt.com/about/our-story/> (last accessed Mar. 22, 2019).

¹²²⁸ *Id.*

¹²²⁹ *Id.*

¹²³⁰ See MALLINCKRODT PHARMACEUTICALS, *Corporate Fact Sheet*, 1 (Mar. 2018), http://www.mallinckrodt.com/globalassets/documents/corporate/corporate-fact-sheet_march-2018.pdf.

worldwide,¹²³¹ two of Mallinckrodt's self-described strengths are being "experts in navigating the often complex terrain that comes with regulatory oversight, particularly concerning the **carefully controlled products** we manufacture," and "acquiring and handling highly regulated and complex raw materials and **controlled substances** [which] give us **unique advantages globally in the opioid** and rare disease arenas ...¹²³²

In December 2018, Mallinckrodt Pharmaceuticals announced a new plan to split into two companies.¹²³³ Specialty Generics would retain the Mallinckrodt name and contain the active pharmaceutical ingredient and opioid manufacturing component of the company.¹²³⁴ The other, Specialty Brands, would focus on innovative specialty pharmaceutical brands.¹²³⁵

The manufacture and sales of controlled substances, including opioid products is a major focus of Mallinckrodt's generic business. In fact, between 1996 and 2017, Mallinckrodt was a leading manufacturer of generic opioid products, selling over \$18 billion in opioid products.¹²³⁶

The Mallinckrodt business model involved the sale of large volumes of generic finished opioid dosage units (finished tablets) to various distributors or wholesalers such as McKesson, Cardinal, and Amerisource Bergen, which in turn, supplied various retail pharmacy customers, both independent pharmacies and retail national chains (e.g., CVS and Walgreens).¹²³⁷ Mallinckrodt was also a direct distributor or wholesaler to various chain customers including CVS and Kroger Co., a retail grocery store chain with embedded pharmacies.

In 2017, Mallinckrodt entered into an agreement with the DEA to resolve the Agency's ongoing investigations and settled allegations made by the DEA that the company prior to 2012 failed to maintain and operate an effective anti-diversion program.¹²³⁸

¹²³¹ *Id.*

¹²³² See MALLINCKRODT PHARMACEUTICALS, *Core Strengths*, <http://mallinckrodt.com/about/core-strengths> (last accessed Mar. 22, 2019).

¹²³³ See Presentation by Mallinckrodt Pharmaceuticals, Planned Spin-off of Specialty Generics Business (Dec. 6, 2018), available at <http://mallinckrodt.com/investors/presentation-documents/>.

¹²³⁴ *Id.* at 4.

¹²³⁵ *Id.*

¹²³⁶ See Jan. 30, 2019 Mallinckrodt Response to Interrogatory No. 33 & Ex. E. ["Jan. 30, 2019 MNK Rog Resp."].

¹²³⁷ Throughout this section, the terms "wholesaler" and "distributor" are used interchangeably. Their meanings are identical and connote an organization that provides opioid products from a manufacturer to a pharmacy for dispensing. In other words, a "middle man" in the classic sense of that word.

¹²³⁸ See Administrative Memorandum of Agreement between U.S. Department of Justice, Drug Enforcement Administration and Mallinckrodt plc. and Mallinckrodt LLC., 3-4 (July 10, 2017) ["Mallinckrodt AMOA"]; see also, John Gillies 30(b)(6) Deposition at 239:8-11 (Feb. 7, 2019) (the formal DEA investigation began in September 2011).

14.2 Executive Summary

Despite having access to voluminous data about its customers' customers (e.g., the retail pharmacies and, in certain jurisdictions, dispensing physicians), Mallinckrodt refused to take appropriate actions against its wholesalers, even when presented with evidence that those wholesalers were engaging in diversionary activities. According to the DEA, from January 2008 through September 2011, while there was an exponential increase in diversion involving oxycodone originating in Florida, which Mallinckrodt claimed they the company "still sold excessive amounts of the most highly abused forms of oxycodone, 30 mg and 15 mg tablets, placing them into a stream of commerce that would result in diversion."¹²³⁹

Although Mallinckrodt claimed it knew nothing about the Florida situation,¹²⁴⁰ Mallinckrodt, through its chargeback data, had access to much of the same information as the DEA regarding these distributors and their diversionary activities.¹²⁴¹ In fact, for those distributors where Mallinckrodt was the sole supplier of opioid products, which included Keysource (for all product types that Mallinckrodt supplied to Keysource)¹²⁴² and Sunrise (for all oxycodone products)¹²⁴³, the company had access to the same information as the DEA.¹²⁴⁴ Rather than cut-off those distributors, which operated in an almost perpetual state of non-compliance with controlled substances requirements, the most Mallinckrodt was willing to do was cut chargeback payments to certain pharmacies, and even then, Mallinckrodt often relented under distributor pressure.¹²⁴⁵

This failure to act on the data in its possession led the DEA in 2017 to accuse Mallinckrodt of not:¹²⁴⁶

- Conducting adequate due diligence of its customers,

¹²³⁹ See AMOA at 1.

¹²⁴⁰ See e.g., email chain between Kate Muhlenkamp-Neely and Victor Borelli re Sunrise license suspensions due to oxycodone sales to Florida (June 21, 2010), MNK-T1_0000383311; Memo from Howard Davis to Karen Harper. (Nov. 4, 2010) (warning that several Florida pharmacies are receiving oxycodone shipments through multiple Mallinckrodt distributors), MNK-T1_0000269410; Email from customer service rep Polly Jordan to Victor Borelli asking if he "heard anything about Oxycodone in Florida, and why there are so many people from Kentucky going to Florida to get their prescriptions filled?" (Mar. 4, 2009), MNK-T1_0000384265.

¹²⁴¹ See Steven Becker Deposition at 312:11-321:12 (Dec. 19, 2018) (acknowledging that Mallinckrodt had same info as DEA re the company's sales in its chargeback system and could have evaluated same data that DEA evaluated, particularly where Mallinckrodt was the sole opioid supplier to a particular downstream customer).

¹²⁴² See Email from Victor Borelli to James Rausch (Mar. 23, 2010), MNK-T1_0000312043 ("Fortunately, on all of our products at [Keysource], we are their sole supplier.")

¹²⁴³ See Email from Victor Borelli to Kate Muhlenkamp et al. (Nov. 12, 2008), MNK-T1_0000565624 ("We have displaced all competitors at this account, and they are relying on our supply to cover their demand.")

¹²⁴⁴ *Id.*

¹²⁴⁵ See "Top 20" spreadsheet for 2011 listing 20 chargeback restricted pharmacies in Florida and another 20 outside of Florida (Sept. 30, 2011), MNK-T1_0000293605. Mallinckrodt approved chargebacks for several pharmacies based on the representations and/or info provided by Cardinal, without requiring an on-site audit first. The spreadsheet contains notes of discussions with Cardinal for each of the 40 pharmacies, for e.g. for three Florida CVS pharmacies, including two in Sanford, the spreadsheet states "take CVS off the list for now, additional dialog (sic) will be conducted with CVS" ... For SASB Inc. dba Okeechobee Discount Drugs, it states "ok to take off the list per agreement between Mallinckrodt & Cardinal, Cardinal will send case file."

¹²⁴⁶ See AMOA at 2.

- Detecting and reporting to the DEA orders of unusual size and frequency, as well as those orders deviating substantially from normal patterns including, but not limited to, those identified in the 2006 and 2007 Rannazzisi letters
- Using ‘chargeback’ information from its distributors to evaluate suspicious orders, and
- Taking effective action to prevent recurrences of diversion by downstream customers after receiving concrete information of diversion of Mallinckrodt’s products by those downstream customers.

In the settlement agreement, Mallinckrodt failed to accept full responsibility for the situation, and simply conceded that “at certain times ... prior to January 1, 2012, certain aspects of Mallinckrodt’s system to monitor and detect suspicious orders did not meet the standards outlined in letters from the DEA Deputy Administrator, Office of Diversion Control” in 2006 and 2007.¹²⁴⁷

However, the DEA’s findings were mere symptoms of a far deeper problem at Mallinckrodt – a failure of culture. Mallinckrodt demonstrated repeatedly that its purported company values of being patient-centric and operating with integrity were just platitudes, and that Mallinckrodt apparently was indifferent to any negative societal impact flowing from its actions. Mallinckrodt’s callousness towards the ultimate consumers of its opioid products was typified in an email exchange between Mallinckrodt’s National Account Manager, Victor Borelli, and one of his distributor contacts at KeySource Medical, Steve Cochrane, after Mr. Cochrane had received an overnight shipment of 1200 bottles (e.g., 120,000 dosage units of oxycodone) from Mallinckrodt:

Mr. Cochrane wrote: “Keep’em comin’! Flyin’ out of here. It’s like people are addicted to these things or something. Oh, wait, people are . . .”

Mr. Borelli responded: “Just like Doritos keep eating. We’ll make more.”¹²⁴⁸

When this email exchange was reported in *The New York Times*, a Mallinckrodt spokesman responded that it was “an outrageously callous email from an individual who has not been employed by the company for many years,” and added that “[i]t is antithetical to everything that Mallinckrodt stands for and has done to combat opioid abuse and misuse.”¹²⁴⁹ However, Mallinckrodt’s current Chief Commercial Officer and previous president of the company’s generics division, repeatedly refused under oath to express any opinions about the behavior, let alone condemn it as being “antithetical” to Mallinckrodt’s corporate values.¹²⁵⁰

¹²⁴⁷ *Id.* at 4.

¹²⁴⁸ See Email chain between Victor Borelli and Steve Cochrane Re Oxy 30 (Jan. 27, 2009), MNK-T1_0000559532.

¹²⁴⁹ See Associated Press, *Suit: US Drug Agency Deemed Firm ‘Kingpin’ in ‘Drug Cartel,’* THE NEW YORK TIMES (Apr. 1, 2019), <https://www.nytimes.com/aponline/2019/04/01/us/ap-us-opioid-lawsuit-tennessee.html>; see also, email from Victor Borelli to Steve Cochrane referenced in the NYT article. (Jan. 27, 2009), MNK-T1_000055953. Note that the “Doritos” conversation occurred over email not via telephone as reported in the article.

¹²⁵⁰ See Hugh O’Neill Deposition at 152:15-153:4; 155:23-158:15; 166:10-169:15; 171:10-175:19 (Mar. 13, 2019).

14.3 Impact

This systemic cultural failure, in turn, resulted in Mallinckrodt's predictable failure to implement a credible and effective anti-diversion program during the review period. Although Mallinckrodt employed a "peculiar order" approach, described as an algorithm plus due diligence on orders deemed to be "peculiar," Mallinckrodt's approach to implementing the "peculiar order" process was utterly ineffective as the table below shows.

Mallinckrodt Suspicious vs. Peculiar Orders 2003-2011

Years	Total Peculiar Orders	Total Suspicious Orders Reported
2003-2007	1,295	10-24
2008-2011	36,522	9

During the same period (2003 to 2011), Mallinckrodt shipped more than 53 million orders of opioid products.¹²⁵¹ Thus, out of 53 million orders only a maximum of 33 apparently never shipped.¹²⁵² Assuming these numbers are accurate, they are indicative of an anti-diversion program was simply *pro forma* exercise.

Mallinckrodt's lack of a credible anti-diversion program also impacted Summit and Cuyahoga Counties. For those counties, Mallinckrodt via one of its subsidiaries was the largest supplier of opioid products. Thus, between 2006-2014, Mallinckrodt shipped more opioid products – 26.7% of the total or 2,363,328,618 MMEs – than any other manufacturer.¹²⁵³

Despite its sizeable market share, Mallinckrodt appears to have expressed zero concern that its opioid products were being diverted in Ohio. Upon receiving a news article describing the closure of a notorious pain clinic in southern Ohio along with the arrest and prosecution of its owner and a doctor at the clinic, Kevin Becker, a Mallinckrodt District Sales Manager, emailed his Regional Sales Director, Jay Meyer the following:

"The good news he was not an Exalgo customer. Bad news is he is aligned to Heidi [a Mallinckrodt sales rep]. I did meet him. Nice guy. Oh well."¹²⁵⁴

This reaction, and complete lack of concern, was not atypical.

Mallinckrodt possessed information regarding problems with diversion in Ohio, as its employees received email alerts from law enforcement regarding investigations and circulated news reports and internal emails regarding

¹²⁵¹ MNK-T1_0007965587 & 5588 (Mallinckrodt sales data).

¹²⁵² *Id.*

¹²⁵³ See Supplemental Expert Report of Craig J. McCann, Ph.D., CFA, ¶ 13 & Table 1 (April 15, 2019).

¹²⁵⁴ Email from K. Becker to J. Meyer, (Dec. 28, 2011) MNK-T1_0005032855.

Ohio pill mills.¹²⁵⁵ Yet Mallinckrodt's former Chief Security Director testified that he was not aware of any particular problem with Ohio.¹²⁵⁶ In short, the opioid situation in Ohio appeared to not be a concern for Mallinckrodt despite the company possessing information demonstrating that would be extremely concerning to a prudent and responsible opioid manufacturer.

Given this, it is unsurprising that Mallinckrodt's distributors shipped opioid products to various Summit and Cuyahoga pharmacies that engaged in questionable activities involving opioids. These included Euclid Family Pharmacy and Walgreens Stores 3226, 3314, and 6574. Mallinckrodt's opioid products were supplied to all these pharmacies.¹²⁵⁷

Moreover, it would be incorrect to limit Mallinckrodt's impact on Cuyahoga and Summit Counties to only those opioids that were directly shipped to these jurisdictions. Prescription opioids cross state lines, and opioids sent to jurisdictions with lax regulation, for example Florida, can easily make their way to Ohio, a fact that Mallinckrodt clearly was aware of.¹²⁵⁸

Responsible compliance departments in possession of this knowledge would have expanded their focus beyond Florida to other jurisdictions in order to ascertain whether there were systemic compliance or diversion problems. There is nothing in the record indicating that Mallinckrodt ever did this for Ohio.

14.4 Company Commitment - Compliance Culture, Organization & Resources

14.4.1 Mallinckrodt's public commitment to its values of being patient-centric and demonstrating integrity does not mirror its actual practice.

Mallinckrodt has publicly stated that it is committed both to the patients who need their opioid products and to conducting their affairs with integrity, including following all laws and regulations. As expressed by Mallinckrodt's Chief Commercial Officer, Hugh O'Neill, "[w]hat we do in our business is always serious business."¹²⁵⁹ However, Mallinckrodt's actions paint a starkly different picture; one in which "serious business" means only the bottom line and not doing the right thing. Simply put, Mallinckrodt does not "walk

¹²⁵⁵ See, e.g., Email from Det. Dennis Luken to rxnews@listserve.com, [RXNews] Forged Prescription Organization, (Oct. 31, 2011) (email from law enforcement regarding organized group of doctors in Cleveland metropolitan area issuing fraudulent prescription for opioids), MNK-T1_0006029397; Email from Collidge to NADDI, [RXNews] FBI news, (Apr 20, 2012) (email regarding indictment of owner of three Ohio pain clinics), MNK-T1_000603138).

¹²⁵⁶ See William Ratliff Deposition at 33:4-14 (Dec. 19, 2018).

¹²⁵⁷ See MNK-T1_0007965587-7965588 (Mallinckrodt chargeback data).

¹²⁵⁸ See Appendix A, Figure 1. Ms. Harper gave a presentation on the "Oxy Express," the shorthand reference for the migration of pills from Florida to Ohio along the I-75 corridor. See Karen Harper Deposition at 91:9-92:19 (Jan. 15, 2019). John Gillies, agreed that it was "common knowledge" that people traveled to Florida to buy opioids because of the state's "lax business practices," and then transported these drugs out of the state. See J. Gillies 30(b)(6) Deposition at 40:8-18. In fact, Steven Becker, one the generic NAMs, was aware of the migration of pills from Florida to other states and the Oxy Express, even as he was shipping millions of pills to Florida. See Steven Becker Deposition, 114:1-18 (Dec. 19, 2018).

¹²⁵⁹ See H. O'Neill Deposition at 164:12-14.

the talk” of corporate and social responsibility, particularly as it relates to the sale and distribution of opioid products, and that tone comes from the top.

A. Code of Conduct

Although Mallinckrodt states that integrity is one of its four values and describes itself as an expert at navigating the regulatory terrain and handling controlled substances,¹²⁶⁰ Mallinckrodt’s 42 page Guide to Business Ethics, subtitled “Integrity is Our Focus,” references controlled substances only twice, and both times in the context of prohibiting employee’s use of or being under the influence of controlled substances in the workplace.¹²⁶¹ Therefore, while Mallinckrodt understood and was concerned over the dangers of opioid misuse in the workplace, the company failed to express a similar corresponding understanding and concern about the societal impact of misusing opioids. This lack of regard for the societal impact is visible in the actions of Mallinckrodt’s sales representatives as discussed below.

B. Sales Representative Words and Actions

The words and actions of the Mallinckrodt sales and marketing department, including statements made by individual sales employees, indicate a singular focus on the sale of opioid products, and an apparent callous disregard for societal norms or for the harm that Mallinckrodt’s opioid products could cause. As noted by Ms. Harper, this callousness towards DEA’s controlled substances requirements was fed by a belief the “we are ‘such big players’ the DEA would never suspend our license.”¹²⁶² A notion, she tried to disabuse at least Mallinckrodt’s compliance employees of.

Given Mallinckrodt’s dependence on its field force to support its lightly resourced Controlled Substances Compliance Group, this is troubling. Furthermore, based on the number of examples uncovered, I do not believe these are isolated actions of a few “rogue sales representatives,” but rather demonstrates a profoundly flawed corporate culture.

It seems that in a misguided attempt to motivate its sales force to higher levels of performance (e.g., more sales) Mallinckrodt developed songs about its opioid products, bearing names such as “Don’t Mess Around,” “Give Me Moore,” “Sweet Relief,” “We Are Covered,” “When Good Ain’t Good Enough,” and “When Less Can Be More.”¹²⁶³ The lyrics of Mallinckrodt’s 2012 song, “Propah Dose,” aptly illustrate the issue.¹²⁶⁴

You can start at the middle

¹²⁶⁰ See Discussion *infra*.

¹²⁶¹ See Mallinckrodt, *Guide to Business Conduct*, 29 (Jan. 8, 2018), http://mallinckrodt.com/globalassets/documents/mallinckrodt_code_external_010818.pdf.

¹²⁶² See Email from K. Harper to B. Ratliff, *et al.*, News on DEA Suspension of License, Amerisource-Bergen, FL, (Apr. 27, 2007) (reminding those sent the email that even big players can be suspended), MNK-T1_0000273471.

¹²⁶³ See announcer script for a Mallinckrodt internal radio show containing lyrics to multiple songs about Mallinckrodt opioid products (including “Don’t Mess Around,” “Give Me Moore,” “Propah Dose,” “Sweet Relief,” “We Are Covered,” “When Good Ain’t Good Enough,” and “When Less Can Be More”), MNK-T1_0002734994.

¹²⁶⁴ Email from M. Falcone to sales personnel announcing the availability of the Propah Dose song. (May 11, 2012), MNK-T1_0002785759.

You can start at the top
You can start with very little
But that's not where you should stop
Cause your patient needs relief, mon
...
So when you start at the middle
Or you start at the top
Or you start with a little
Make sure you don't stop
Cause your patient needs relief, mon¹²⁶⁵

It, however, did not stop there as Mallinckrodt's sales department commissioned a video, apparently modeled on the "Dos Equis" beer commercial, which described "the most interesting physician in the world," and intoned that "[e]very time she writes a prescription, an angel gets its wings."¹²⁶⁶ The fact that a portion of Mallinckrodt's sales and marketing budget was devoted to these efforts is indicative of senior leadership indifference, but it also reflects a poor "tone at the top" that is not committed to compliance.

This poor compliance culture ultimately permeated the entire organization resulting in individual employees exercising poor judgment and engaging in unbecoming behavior. Perhaps the best example of how Mallinckrodt's poor compliance culture negatively impacted its employees' behavior can be seen with Victor Borelli, one of Mallinckrodt's most highly compensated National Account Managers ("NAMs").

Mr. Borelli worked for Mallinckrodt from 2005 to 2012. Before joining Mallinckrodt, Mr. Borelli was the director of coffee products for Sara Lee, and thus did not have any experience with either the pharmaceutical industry or Schedule II narcotics. In fact, Mr. Borelli was unable to identify whether certain of the products he was responsible for selling were even opioids.¹²⁶⁷ Nor could he recall the specifics of any compliance training.¹²⁶⁸

However, as his compensation records show, Mr. Borelli was quite successful at developing relationships with his accounts.¹²⁶⁹ However, Mr. Borelli's callousness and lack of concern for his compliance responsibilities, as demonstrated by the "Doritos" email¹²⁷⁰ detailed above and other conduct, was a cause for concern among other staff members with controlled substance responsibilities. For example, Customer Service Manager Cathy Stewart warned Ms. Harper and Mr. Ratliff that the customer service representatives all said that "Borelli will tell them anything they want to hear just so he can get the sale."¹²⁷¹ Mr. Borelli even characterized his job at Mallinckrodt using the phrase "ship, ship, ship."¹²⁷² Finally, in a blatant disregard for anti-diversion

¹²⁶⁵ See Propah Dose lyrics, MNK-T1_0002734994 at 2375001.

¹²⁶⁶ See Dos Equis spoof ad, MNK-T1_0007033463.

¹²⁶⁷ See Victor Borelli Deposition at 55:19-56:19 (Nov. 29, 2018).

¹²⁶⁸ *Id.* at 28:9-29:23.

¹²⁶⁹ Excel sheet showing national account managers and bonus payments for 2007-2011, MNK-T1_0000315995.

¹²⁷⁰ Email chain between Victor Borelli and Steve Cochrane Re Oxy 30 (Jan. 27, 2009), MNK-T1_000559532.

¹²⁷¹ See Email from Cathy Stewart to Bill Ratliff and Karen Harper Re Sunrise Wholesale (May 20, 2008), MNK-T1_0000290611.

¹²⁷² See Email chain between Victor Borelli and Kate Muhlenkamp-Neely (Sept. 16, 2008), MNK-T1_0000563696.

compliance, Mr. Borelli emailed a distributor client asking them to check their inventories and “[i]f you are low, order more. If you are okay, order a little more. Capesce?”; and joked that the distributor should “destroy this e mail... Is that really possible? Oh Well...”¹²⁷³

Mr. Borelli, however, was not a “rogue sales person,” but a product of Mallinckrodt’s poor compliance environment set from the top. Hugh O’Neill, previously the president of Mallinckrodt’s generic division before becoming the current Chief Commercial Officer, when confronted with a number of instances of troubling behavior or conduct by the Mallinckrodt sales department, including Mr. Borelli’s, consistently and repeatedly testified that he was unable to express any opinions about the behavior, let alone condemn it as being inconsistent with Mallinckrodt’s corporate focus on “integrity” and compliance.¹²⁷⁴ Consequently, a member of Mallinckrodt’s senior leadership claimed not to understand his company’s espoused values or to be able to make a relatively “black or white” integrity call. This is an example of poor ethical leadership and company culture.

14.4.2 Mallinckrodt failed to appropriately organize and staff its anti-diversion program rendering it ineffective.

Throughout the review period, Mallinckrodt’s organization and staffing of the its anti-diversion compliance team, as well as its reliance on its sales force to support the program revealed a fundamental lack of appreciation about the nature, role and importance of the anti-diversion and suspicious order monitoring programs to both Mallinckrodt and the patients it serves.

A. Organizing and Staffing the Controlled Substances Compliance Group

Prior to March 2008, there is scant evidence that Mallinckrodt had a formally designated SOM team.¹²⁷⁵ Mallinckrodt’s anti-diversion compliance team was named first, the DEA Compliance Group (“DCG”) and later, the Controlled Substances Compliance (“CSC”) Group.¹²⁷⁶ However, Mallinckrodt’s actions in organizing and staffing the CSC Group demonstrate that its public statements concerning the importance of controlled substances compliance withstanding, Mallinckrodt really did not value the program. For example, Mr. O’Neill, could not even accurately describe Karen Harper’s role at the company (even though she had worked there for 40 years).¹²⁷⁷ Ms. Harper currently serves as Mallinckrodt’s Director of Controlled Substances Compliance.¹²⁷⁸

¹²⁷³ See Email from Victor Borelli to Steve Cochran, Re. things (May 20, 2008), MNK-T1_0000506535.

¹²⁷⁴ See H. O’Neill Deposition at 152:15-153:4; 155:23-158:15; 166:10-169:15; 171:10-175:19.

¹²⁷⁵ See Mallinckrodt, *Suspicious Order Monitoring Team Charter* (Apr. 7, 2011), MNK-T1_000496062 (showing team start date as 03/28/08); see also, J. Gillies 30(b)(6) Deposition at 101: 7-102:2 (admitting to not seeing any documentation of a SOM team or SOM program before March 2008). Mr. Gillies currently is Vice President of Global Security for Mallinckrodt.

¹²⁷⁶ See K. Harper Deposition at 23:11-17; 30:11-12.

¹²⁷⁷ See H. O’Neill Deposition at 153:17-154:9 (testifying that Harper “was part of the financial organization” and that he could not recall if she had any responsibilities with respect to DEA compliance and suspicious order monitoring.).

¹²⁷⁸ See K. Harper Deposition at 30:11-12.

Also, regardless of the group's name, it was small, consisting of anywhere between 3-8 individuals from 2003 to the present.¹²⁷⁹ As a result, for a two-year period from 2008 through 2010, a single employee – James Rausch – was tasked with reviewing all the peculiar order reports, even though this was not his full-time job.¹²⁸⁰ Consequently, during this period Mr. Rausch resorted to releasing peculiar orders prior to completing his due diligence on them because he didn't have enough time to complete his work.¹²⁸¹

The CSC resource limitation was further compounded by the fact that its key personnel lacked specific controlled substances experience, expertise, and training. For example, Karen Harper, who currently is the Director of Controlled Substance Compliance and whom others in Mallinckrodt considered as the definitive source for questions about the company's SOM and anti-diversion programs,¹²⁸² had no controlled substances experience prior to joining the group, and her only training after she became a senior manager in these areas consisted primarily of attending one industry training conference per year.¹²⁸³

B. Using the Mallinckrodt Sales Force

To supplement the limited CSC staff, Mallinckrodt relied on its sales personnel including its NAMs and Customer Service Representatives ("CSRs") to serve in active anti-diversion roles. The NAMs were considered the "eyes and ears and boots on the ground at the customer accounts," chiefly assisting in the vetting of peculiar orders while the CSRs were "veteran[s] in the business, and ... in general familiar with customer's order patterns."¹²⁸⁴

Despite the belief that the NAMs were trained to "be vigilant for any potential sign – red flags that could be indicative of diversion as they visited customers,"¹²⁸⁵ Mallinckrodt overlooked the fact that the compensation structure for the NAMs was weighted heavily to favor sales over compliance. For example, one NAM's bonuses (Victor Borelli) nearly tripled from \$35,000 in 2007 to more than \$100,000 in 2008-2010.¹²⁸⁶ The NAM compensation plan was so lucrative that Mr. Borelli exceeded his sales goals with one of his top customers, Masters Pharmacy, by more than 1,400%.¹²⁸⁷

While compensation plans based on product sales are not *per se* wrong, in Mallinckrodt's case because there were no counterbalancing incentives for reporting suspicious customer orders and activity, or performing the

¹²⁷⁹ See K. Harper Deposition at 24:23-25:2; 27:2-7.

¹²⁸⁰ See James Rausch Deposition at 133:18-134:8; 213:13-215:24 (Nov. 15, 2018).

¹²⁸¹ See Email from James Rausch to Karen Harper (June 9, 2010), MNK-T1_0000279153; *see also*, James Rausch Deposition at 112:2-114:7 (Nov. 16, 2018) (noting that Rausch had an "agreement" with Harper that peculiar orders could be shipped prior to due diligence being completed because Mallinckrodt "didn't always have the ability to do the thorough investigation prior to the order being shipped."); *see also* Discussion *infra*.

¹²⁸² See W. Ratliff Deposition at 146:8-19 (testifying that he believed Harper was "very informed and knows everything there is about DEA compliance"); *see also*, J. Rausch Deposition at 129:8-12; Ginger Collier Deposition at 214:11-24 (Jan. 8, 2019).

¹²⁸³ See K. Harper Deposition at 24:8-12; 49:17-50:25.

¹²⁸⁴ See K. Harper Deposition at 59:15-16; 59:23-25.

¹²⁸⁵ *Id.* at 59:16-19.

¹²⁸⁶ Excel spreadsheet showing National Account Manager bonus payments, MNK-T1_0000315995.

¹²⁸⁷ Excel sheet showing 2008 net sales and net sales objectives, MNK-T1_0000562520. (Borelli Deposition Ex. 4).

other anti-diversion tasks assigned to them, the company put a system in place that clearly valued sales and revenue above compliance and corporate responsibility. As a result, the NAMs were viewed as “advocates” for their distributor customers.¹²⁸⁸ Furthermore, Ms. Harper knew that as of November 2008, it would not be a good idea to consult with NAMS in connection with the approval of new customers, and that sales people should not be involved heavily in the identification of suspicious orders.¹²⁸⁹

While the NAMs’ bonuses were tied to their generic opioid sales, and those bonuses could reach six figures,¹²⁹⁰ Mallinckrodt apparently did not evaluate or compensate NAMs on their compliance responsibilities. Nor is there anything in the record indicating that a NAM was ever penalized for failing to stop a suspicious order. Kate Neely (formerly Muhlenkamp), the product manager for oxycodone for three years and who helped coordinate the SOM process, testified that she could not recall a single instance in which she recommended blocking a shipment of opioids to the CSC Group.¹²⁹¹

Mallinckrodt also had received warnings about its lopsided compensation system from its own staff. In 2008, Mallinckrodt’s compliance personnel attended an industry conference where they learned that it was not advisable to rely so heavily on sales personnel in the SOM process.¹²⁹² Two years later, September 2010, Ms. Harper recommended that “the actual day-to-day monitoring responsibility should be switched to a non-customer service function in that those that have responsibility to manage the orders have a conflict of interest in deciding which orders should ultimately be shipped – with ultimate right of refusal retained by Controlled Substance Compliance.”¹²⁹³ However, despite these warnings, Mallinckrodt did not address the conflict situation.

14.5 Program Core – Requirements, Education, Detection & Corrections

14.5.1 Mallinckrodt’s poor documentation practices were an impediment to the company’s efforts to establish an effective diversion program.

Throughout the review period, Mallinckrodt’s anti-diversion program, including suspicious order monitoring, was hamstrung by the company’s lack of adherence to good documentation practices surrounding the development, deployment and use of written standards. Consequently, this failure undermined the overall effectiveness of the controlled substances program, and it did not comport with the compliance program standards and industry leading practices.

¹²⁸⁸ See Kate Neely Deposition at 342:5-8 (Jan. 8, 2019).

¹²⁸⁹ See K. Harper Deposition at 283:10-288:9; see also K. Harper Exhibit 13 (Nov. 14, 2008 email from Cathy Stewart attaching notes from the DEA Buzzeo conference, MNK-T1_000043222-226).

¹²⁹⁰ Excel spreadsheet showing National Account Manager bonus payments, MNK-T1_0000315995.

¹²⁹¹ See K. Neely Deposition at 354:21-355:21.

¹²⁹² See Meeting Notes from Buzzeo Conference (Oct. 27-30, 2008), MNK-T1_0000302097 (reporting the “general consensus is that sales reps are not considered a good option for on-site investigations and initial review prior to accepting new customers due to their perceived bias in getting the customer approved for sales revenue purposes.”).

¹²⁹³ See Email from Karen Harper to Eileen Spaulding Re SOM (Sept. 24, 2010), MNK-T1_0000280260.

Prior to 2008, Mallinckrodt's anti-diversion program was marked by an absence of formal written standards. Although Karen Harper, currently Mallinckrodt's Director of Controlled Substance Compliance, noted that Mallinckrodt had a SOM program since at least 2003,¹²⁹⁴ it was not until 2008 that documentation in the form of a draft SOP made an appearance.¹²⁹⁵ She also described the program during that period as consisting of an "algorithm and some other factors."¹²⁹⁶

The written standards for suspicious order monitoring and anti-diversion in effect from 2008 to 2015 are best described as chaotic. This period was marked by intensive activity that resulted in the creation of a complex and confusing set of written standards. Consequently, Mallinckrodt's anti-diversion program in this period lacked both clarity and consistency.

Ms. Harper and the CSC group began work on a suspicious order monitoring SOP sometime in April or May 2008.¹²⁹⁷ In October 29, 2010, a "finished" version of the SOM policy was created.¹²⁹⁸ While it appears to be a final version of the SOP, Ms. Harper acknowledged that as of October 31, 2010 "a final SOM procedure that outlines the criteria for identifying a suspicious order had not been finalized."¹²⁹⁹ Therefore, although work started in 2008, the finished SOP still was not completed and finalized at the end of October 2010, more than two years later.

According to Ms. Harper, it also was Mallinckrodt's practice that any operative drafts that CSC was "working on at the time was also the policy that [the company] would follow with respect to Mallinckrodt's suspicious order monitoring obligations."¹³⁰⁰ Therefore, under this process, the drafts were essentially "approved" policies or procedures, but "approved" informally only by Ms. Harper and a small group of colleagues. Thus, it appears that Mallinckrodt did not have, or Ms. Harper did not utilize, an appropriate formal document control process, making it virtually impossible to discern what the final established policy was for the SOM program at any particular point in time.

This difficulty was further compounded by that fact that beginning with the October 2010 version of C/S 3.0, Mallinckrodt ceased marking operative drafts as "drafts." Prior to this version, Ms. Harper had marked the various 2008 versions as "published drafts."¹³⁰¹

¹²⁹⁴ See K. Harper Deposition at 83:17-23 (confirming that the first reference to the existence of a SOM program at Mallinckrodt was 2003). Ms. Harper's testimony is particularly relevant because she was identified, and she confirmed, that given her long tenure with Mallinckrodt, she was most knowledgeable about the history of Mallinckrodt's anti-diversion program. See *id.* at 70:11-71:10.

¹²⁹⁵ See, e.g., Mallinckrodt DEA Compliance Procedure, *Controlled Substance Suspicious Order Monitoring*, Draft 2 (May 13, 2008), MNK-T1_0000268911.

¹²⁹⁶ See K. Harper Deposition at 83:24-84:3.

¹²⁹⁷ See Mallinckrodt DEA Compliance Procedure, *Controlled Substance Suspicious Order Monitoring*, Draft 2 (May 13, 2008), MNK-T1_0000268911.

¹²⁹⁸ See Mallinckrodt Global Controlled Substances Compliance Procedure, *Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program, C/S Comp. 3.0* (Oct. 29, 2010), MNK-T1_0000264260.

¹²⁹⁹ See K. Harper Deposition at 323:11.

¹³⁰⁰ See K. Harper Deposition at 435:14-20.

¹³⁰¹ See, e.g., Mallinckrodt DEA Compliance Procedure, *Controlled Substance Suspicious Order Monitoring*, Draft 2 (May 13, 2008), MNK-T1_0000268911; see also Appendix H, Figure 1 *infra* (listing the other versions of this document).

The entire approach of publishing and using operative drafts is an example of exceedingly poor document control practices and corporate governance. Drafts should never be used as “final documents,” because by their very nature, they are subject to further and sometimes frequent revision. The history of Mallinckrodt’s SOM procedure aptly demonstrates this. From 2008 to 2015, the *Controlled Substances Suspicious Order Monitoring* procedure and its progeny were modified fifteen times, and in the case of 2008, 2011, and 2012, there were three or more revisions per year.¹³⁰² Nor was it until March 2011 that a document “revision history” section for the compliance written standards made its first appearance.¹³⁰³ Thus, this need for further and frequent revisions undermined the primary purpose of written standards, namely, to provide operational clarity and consistency.

Although it is not unexpected for compliance programs to have some policies and procedures that are “off-line” because they are new or being revised to reflect current operational practices, normally these periods are short-lived. When these situations occur, the organization committed to compliance will recognize the gaps or inconsistencies and create a plan to remedy them in a timely manner (e.g., a corrective action plan). However, it is outside the norms of good corporate governance, as well as a symptom of an organization’s poor adherence to basic document control standards, for an organization to allow “drafts” containing gaps and inconsistencies to be used in place of final written standards and to do so unabated for periods of years, like Mallinckrodt did in this situation.

This informal approach to written standards delivered a predictable outcome; most notably confusion. For example, in 2012, while searching for Mallinckrodt’s policy regarding the identification/investigation of suspicious orders, Gail Tetzlaff, Mallinckrodt’s Director of Government Affairs, wrote: “I would assume this is going to be the first official Revision but when I looked in the SOP folder on the Shared Drive, I didn’t see a finalized copy of our original version. What is the status of that?”¹³⁰⁴

The lack of formalized written standards outlining Mallinckrodt’s suspicious order monitoring program was a major shortcoming that undermined Mallinckrodt’s claims about the company’s efforts to meet its anti-diversion obligations. It also was not in sync with either corporate compliance program standards or industry practices.¹³⁰⁵ Furthermore, the need for formal written standards was something that Mallinckrodt knew or should have known it needed.¹³⁰⁶

¹³⁰² See Appendix H, Figure 1, *infra*.

¹³⁰³ See Mallinckrodt Global Controlled Substances Compliance Procedure, *Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program, C/S Comp. 2.0*, (Mar. 28, 2011), MNK-T1_0000264209; *but cf.*, Mallinckrodt Global Controlled Substances Compliance Procedure, *Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program, C/S Comp. 3.0* (Oct. 29, 2010), MNK-T1_0000264260.

¹³⁰⁴ Email from Gail Tetzlaff to Jennifer Buist, Re draft changes to SOM procedure (Oct. 8, 2012), MNK-T1_0008246445.

¹³⁰⁵ See Discussion *infra*.

¹³⁰⁶ See generally Discussion *infra* at Sections 4 and 5.

14.5.2 Mallinckrodt used the artifice of “peculiar orders” to avoid reporting suspicious orders to the DEA.

The concept of “peculiar orders” seems to first appear in early 2008 with Ms. Harper’s initial draft of the controlled substances suspicious order monitoring procedure.¹³⁰⁷ As defined in that draft, a peculiar order was “[a] controlled substance order that meets an internal established criteria that will not be shipped pending further review by DEA Compliance.”¹³⁰⁸ On the other hand, a suspicious order was defined as “[a] controlled substance Peculiar Order that has been reviewed by DEA Compliance and Security that will not be shipped and will be reported to the Drug Enforcement Administration.”

Thus, within the Mallinckrodt program there were three types of orders: regular orders, peculiar orders and suspicious orders. However, according to the DEA’s requirements, there are only two types of orders: regular and suspicious. Mallinckrodt, however, persisted in the belief that “a peculiar order ... is not necessarily synonymous with a suspicious order” because within Mallinckrodt “certain people make the determination of whether or not the order is ultimately suspicious **sufficient to notify the DEA.**”¹³⁰⁹

While the term “peculiar order” is unique to Mallinckrodt, the concept is not. As discussed elsewhere in this report, every distributor at one time within the review period utilized a term of art to describe orders of unusual size, frequency or pattern that was in their opinion not a “suspicious order.”¹³¹⁰ The most common term being “orders of interest.” Therefore, a “peculiar order” and “an order of interest” are one and the same.¹³¹¹ Furthermore, as previously discussed, neither term has any regulatory validity. Moreover, it was not appropriate for Mallinckrodt to deem a “peculiar order” as not suspicious and simply not investigate them further.¹³¹²

Throughout the review period, the definition of “peculiar orders” morphed over time until it finally was removed from the SOM SOP in 2012 to be replaced by the term “suspicious order.”¹³¹³ However, during the time the term was used, the various versions of “peculiar order” remained out of sync with the DEA’s definitions and its expectations.

The first major change to the definition of a “peculiar order” occurred in June 2008, when Mallinckrodt removed the statement that peculiar orders “will not **be shipped** pending further review by DEA Compliance” and replaced it with language that peculiar orders “will be **placed on hold** pending review by DEA

¹³⁰⁷ See Mallinckrodt DEA Compliance, *Controlled Substance Suspicious Order Monitoring*, Revision 1 (undated), MNK-T1_0000273894. However, as Ms. Harper testified, “at different times with the enhancements of our program we [Mallinckrodt] called orders “peculiar,” we called orders “unusual,” and we called orders “suspicious.” See K. Harper Deposition at 190:11-14.

¹³⁰⁸ See K. Harper Deposition at 190:11-14.

¹³⁰⁹ See K. Harper Deposition at 190:11-23; 191:2-9.

¹³¹⁰ See Discussion *infra*.

¹³¹¹ In addition, the words “peculiar” and “suspicious” have almost the same meaning. “Peculiar” means “different to what is normal or expected,” while “suspicious” means “causing one to have the idea or impression that ... something is questionable” See OXFORD ENGLISH DICTIONARY, “PECULIAR”, <https://en.oxforddictionaries.com/definition/peculiar> (last visited Mar. 30, 2019); OXFORD ENGLISH DICTIONARY, “SUSPICIOUS”, <https://en.oxforddictionaries.com/definition/suspicious> (last visited Mar. 30, 2019).

¹³¹² See Discussion *infra* Section 5.4.

¹³¹³ See Covidien, *HZQS – Identification, Investigation, and Reports of Controlled Substances Suspicious Orders*, 1 (Oct. 18, 2012), MNK-T1_0007476261.

Compliance”¹³¹⁴ The change in the definition is consistent with Ms. Harper’s statement that “[t]here were times that we shipped an order before the [due diligence] review was complete, **but we never shipped a suspicious order.**”¹³¹⁵

The fact that Mallinckrodt allowed orders to ship without the further investigation (a.k.a. due diligence) being completed is not in conformance with DEA expectations and industry standards.¹³¹⁶ As Ms. Harper conceded, if a peculiar order was released without doing further investigation or due diligence a potentially suspicious order could be shipped.¹³¹⁷ Thus, Mallinckrodt knew it was shipping orders that potentially were suspicious, but did not stop shipment of them.

With the creation of a more formal SOP in October 2010, Mallinckrodt made significant changes to the definition of “peculiar order.”¹³¹⁸ According to this new SOP version, a peculiar order was a “[c]ontrolled substance order that meets internal established criteria of 3X the average amount of product ordered during the previous 12 months by DEA reporting class.”¹³¹⁹ While the new definition added the level at which an order was “flagged” as peculiar, it removed any reference that “flagged” orders were automatically placed on hold.¹³²⁰

The October 2010 definition remained in place until the entire procedure was replaced in October 2012.¹³²¹ With the new SOP in October 2012, the term “peculiar order” simply became a “suspicious order.” In this version, Mallinckrodt defined a suspicious order as

An order received by Mallinckrodt directly from a customer for a Schedule II – V Controlled Substance product which exceeds the internal limit set by the application of the two-tiered system of specifically created algorithms defined in this policy. Suspicious Orders will be reported to DEA upon discovery in accordance with 21 C.F.R. § 1301.74(b).¹³²²

¹³¹⁴ See Mallinckrodt DEA Compliance, *Controlled Substance Suspicious Order Monitoring*, Revision 1 (undated) (emphasis added), MNK-T1_0000273894; Mallinckrodt DEA Compliance, *Controlled Substance Suspicious Order Monitoring*, Draft #3 (June 2, 2008) (emphasis added), MNK-T1_000419993.

¹³¹⁵ See K. Harper Deposition at 201:17-20.

¹³¹⁶ See, e.g., Letter from Joseph Rannazzisi to All Registrants at 2 (Dec. 7, 2007), MNK-T1 0000263492 (“Registrants who routinely report suspicious orders yet fill these orders without first ascertaining that the order will not be diverted into other than legitimate medical, scientific, or industrial channels, are failing to maintain effective controls against diversion.”); see also HDMA *Position Statement and Industry Compliance Guidelines: Report Suspicious Orders and Preventing Diversion of Controlled Substances* (2008), WAGMDL00673706-673722.

¹³¹⁷ See K. Harper Deposition at 203:1-17.

¹³¹⁸ See Mallinckrodt Global Controlled Substances Compliance Procedure, *Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program*, C/S Comp. 3.0, 2 (Oct. 29, 2010), MNK-T1_0000264260; see also Appendix H, Figure 2 *infra*.

¹³¹⁹ See *id.* at MNK-T1-0000264260.

¹³²⁰ *Id.*

¹³²¹ See Covidien, *HZQS – Identification, Investigation, and Reports of Controlled Substances Suspicious Orders*, 1 (Oct. 18, 2012), MNK-T1_0007476261.

¹³²² *Id.* at 1.

Therefore, 2012 is the first time that Mallinckrodt's definition of suspicious orders began to track with the DEA's definition. However, nowhere in the procedure was there a clear definition that a suspicious order is an order of unusual size, frequency or pattern, but instead it is couched in terms of exceeding an established threshold. Furthermore, there was no precise designation of when a suspicious order is "discovered" (e.g., is it when the algorithm flags the order or when the SOM Leadership team completes its review).¹³²³ The new procedure did reinstate language about automatically generating a "ship-hold" on orders until a final, written determination was made.¹³²⁴ This approach to suspicious orders carried over into the post-2013 era as seen in the 2015 version.¹³²⁵

14.5.3 The Mallinckrodt suspicious order program was flawed both in its design and implementation rendering it ineffective to detect suspicious orders.

Using the concept of "peculiar orders" Mallinckrodt developed a two-tier approach to suspicious order monitoring. The first tier consisted of an automated algorithm to "flag" orders as peculiar. The second tier involved conducting due diligence on the orders identified as peculiar to determine whether they were truly "suspicious" and thus reportable to the DEA.

A. The Peculiar Order Algorithm and Due Diligence

It appears that by 2003 Mallinckrodt employed some sort of automated algorithm to identify suspicious orders.¹³²⁶ Although the algorithm was the primary method for identifying peculiar orders, in some cases both the National Account Managers and Customer Service Representatives could identify an order as "peculiar" independent of the algorithm.¹³²⁷

From March 2008 onwards with the formation of the SOM Team, Mallinckrodt worked to refine and document its algorithmic model.¹³²⁸ That work continued through 2009 and stretched into 2010.¹³²⁹ From its inception, as recounted by James Rausch, Mallinckrodt's Customer Service Manager for Finished Goods, the peculiar order

¹³²³ *Id.* at 2, § 6.4.1 and 3, § 6.4.8.

¹³²⁴ *Id.* at 2, § 6.3.1.

¹³²⁵ See generally, Mallinckrodt Pharmaceuticals, *HZQS – Identification, Investigation, and Reports of Controlled Substances Suspicious Orders*, (Aug. 17, 2015), MNK-T1_0000511246.

¹³²⁶ See K. Harper Deposition at 83:24-84:3 (The Mallinckrodt SOM program at that time employed an "algorithm and some other factors."). However, since there was no dedicated SOM team until March 2008 there is little documentation to support Mallinckrodt's practices in the pre-2008 timeframe. See Mallinckrodt, *Suspicious Order Monitoring Team Charter*, (Apr. 7, 2011) (Showing team start date as 03/28/08), MNK-T1_000496062; see also J. Gillies 30(b)(6) Deposition at 101:7-102:2 (admitting to not seeing any documentation of a SOM team or SOM program before March 2008).

¹³²⁷ See K. Harper Deposition at 196:16-197:10; 197:15-22.

¹³²⁸ See, e.g., Mallinckrodt, *Suspicious Order Monitoring Program Dosage Products Shipments from Hobart – Activities 10/2008 through 8/10*, 1 (undated), MNK-T1_0000477889 ("SOM Team Activity Log").

¹³²⁹ *Id.* at 4 (showing continued interactions between the SOM Team and IT concerning the algorithm).

algorithm was a simple size algorithm that was baked into Mallinckrodt's JD Edwards order entry system¹³³⁰ and flagged orders in excess of [REDACTED]¹³³¹ Orders identified as "peculiar" were compiled into the "Peculiar Order Daily Reports," which were "algorithm reports that show[ed] any orders regarding the unusual size, pattern, et cetera."¹³³²

Orders listed in the Daily Reports were then "reviewed by a customer service representative."¹³³³ However, the review of orders by Customer Service was limited only to those orders on the daily report.¹³³⁴ Therefore, orders that did not trigger the "peculiar order" threshold were not subjected to further scrutiny, even if there may have been other reasons for the order to be deemed suspicious, notwithstanding its relative size.¹³³⁵ Thus, it was possible for suspicious orders not flagged by the algorithm to escape detection and be shipped to the customer.¹³³⁶

Even using the "Peculiar Order Daily Report" and review by a customer service representative, Mallinckrodt could, and at times did, still ship orders on the report. As Mr. Rausch explained to Ms. Harper "[a]s we discussed I do not hold **any** orders while I do my research due to time constraints ... it takes time to get information back from Marketing and sales, sometimes all day."¹³³⁷ He also noted that "[s]ince I don't hold the orders up during my due diligence it's possible that the order could ship. I thought we discussed this and alerting the DEA about a suspicious order/customer for further investigation would be our process going forward."¹³³⁸

Mallinckrodt shipped peculiar orders prior to completing any further due diligence, because it "didn't always have the ability to do the thorough investigation prior to the order being shipped" due to time constraints.¹³³⁹ As Ms. Harper characterized it, "[t]here were times that we [Mallinckrodt] shipped an order before the [due diligence] review was complete, **but we never shipped a suspicious order.**"¹³⁴⁰ During his testimony, Mr. Rausch stated that the company felt comfortable with this practice "because that order was going to a distributor

¹³³⁰ See J. Rausch Deposition at 194:3-10 (the algorithm was part of the computer code for the order entry system, known as JD Edwards or JDE, an Enterprise Resource Planning ("ERP") software platform).

¹³³¹ See J. Rausch Deposition at 44:12-45:16.

¹³³² See J. Gillies 30(b)(6) Deposition at 83:18-22.

¹³³³ See Mallinckrodt DEA Compliance Procedure, *Controlled Substance Suspicious Order Monitoring*, Draft #2, 2 (May 13, 2008), MNK-T1_0000268911.

¹³³⁴ See J. Gillies 30(b)(6) Deposition at 92:7-18 (confirming that Customer Service only reviewed peculiar orders and not every customer order).

¹³³⁵ See J. Rausch Deposition at 194:3-23 (confirming that if an order was not flagged by Mallinckrodt's algorithm, it was not examined; and acknowledging that if there were gaps or faults in the algorithm, it was possible for problematic orders to get through.)

¹³³⁶ *Id.*

¹³³⁷ Email from James Rausch to Karen Harper and George Saffold Re Suspicious Order Monitoring Program (June 9, 2010) (emphasis added), MNK-T1_0000279153.

¹³³⁸ *Id.*

¹³³⁹ See J. Rausch Deposition at 112:14-24.

¹³⁴⁰ See K. Harper Deposition at 201:17-20 (emphasis added); see also Email from K. Harper to J. Rausch and G. Saffold Re Suspicious Order Monitoring Program (June 9, 2010) ("I merely wanted to confirm that we have not shipped any suspicious orders (even if we do the investigation after the shipment)."), MNK-T1_0000279153.

who also had a program in place for suspicious orders, so we knew **we could get the product back or stop it if need be.**¹³⁴¹

How frequently shipments went out without completing due diligence is a matter of some debate. Cathy Stewart, a Customer Service Manager for Mallinckrodt, recounted that she could not recall a single peculiar order being held.¹³⁴² Regardless of how frequently this situation occurred, it was clear that if a peculiar order was released without doing further investigation or due diligence, a potentially suspicious order could be shipped.¹³⁴³

Even when an “investigation” of a peculiar order was undertaken by the Customer Service and the CSC group, it was at best rudimentary, and primarily based upon information provided by the NAMs, who had a conflict of interest. The record indicates that the NAMs simply collected information from their distributor customers and then passed it onto the compliance group, which then simply relied on that information.¹³⁴⁴ In fact, Mr. Borelli simply forwarded inquiries from his compliance department to his wholesale customer, though he claimed to also have followed up with a phone call.¹³⁴⁵

As a result, the justifications used to push orders through generally consisted of two- or three-line emails citing various “marketing rationales.” For example, there was: (a) the fact that a customer was an “established customer”¹³⁴⁶; (b) that the customer had a new account that was ordering a lot of oxycodone¹³⁴⁷; (c) that the customer had expanded its customer base to other states¹³⁴⁸; (d) that the NAM wanted to keep the “momentum rolling” with a customer¹³⁴⁹; (e) that the customer needed to increase its order volume in order to comply with contractual provisions requiring it to order more opioids to secure favorable pricing¹³⁵⁰; (f) that the customer was dramatically increasing its orders to meet customer demand¹³⁵¹; and (g) assuming that a customer’s inability to obtain opioids from another source justified filling an order many times above historical levels.¹³⁵²

¹³⁴¹ See J. Rausch Deposition at 113:23-114:7 (emphasis added).

¹³⁴² See Cathy Stewart Deposition, 111:8-12 (Dec. 11, 2018).

¹³⁴³ See K. Harper Deposition at 203:1-17.

¹³⁴⁴ S. Becker Deposition at 281:13-21; 287:21-13.

¹³⁴⁵ V. Borelli Deposition at 280:2-281:2.

¹³⁴⁶ Email from Bill Ratliff to James Rausch *et al.* Re Propoxyphene Napsylate orders for Sovereign (Sept 3, 2008), MNK-T1_0000290520.

¹³⁴⁷ Email from Steven Becker to Karen Harper *et al.* Re Old Bridge account (Oct. 28, 2008), MNK-T1_0000448888.

¹³⁴⁸ Email from Victor Borelli to Kate Muhlenkamp (Nov. 14, 2008), MNK-T1_0000290502 (noting that new Sunrise Wholesale sales manager is “extremely tied into the Florida market”).

¹³⁴⁹ Email from Victor Borelli to James Rausch Re Peculiar order report (Mar. 23, 2010), MNK-T1_0000297371.

¹³⁵⁰ Emails from Steven Becker to Penny Myers and James Rausch Re HD Smith (May 5, 2010), MNK-T1_0000298906.

¹³⁵¹ Email from Victor Borelli to Brenda Rehkop *et al.* (Jun. 23, 2010), MNK-T1_0000560555 (explaining Oxy’s historical movement through wholesaler and distributor accounts).

¹³⁵² Email from Bill Ratliff to Kate Muhlenkamp *et al.* Re Oxycodone orders (Jun. 4, 2008), MNK-T1_0000562682.

Moreover, if the NAM justified a peculiar order based simply on a distributor's increased sales or market share, the CSC group approved the order on that basis. As Ms. Harper stated in a July 2010 email, "[i]ncreased market share . . . is a satisfactory explanation from the Suspicious Order Monitoring perspective."¹³⁵³

Overall, Mallinckrodt's undue reliance on the peculiar order algorithm, and its shipment of orders flagged by that algorithm with little or no further due diligence is problematic in two significant respects. First, as the DEA counselled, algorithms are not perfect and "[r]egistrants that rely on rigid formulas to define whether an order is suspicious may be failing to detect suspicious orders."¹³⁵⁴ This was known and acknowledged by both Mr. Rausch and Ms. Harper.¹³⁵⁵ It also was something that the customer service department warned CSC about in 2009 noting that the algorithm potentially allowed customers to evade the system by gradually increasing their order volume to avoid triggering the threshold.¹³⁵⁶

Second, there was a DEA expectation that if an order is flagged as suspicious, the company should not ship it until an appropriate investigation is undertaken.¹³⁵⁷ In April 2008, Paul "Pete" Kleissie, DEA Diversion Group Supervisor, inquired about why Mallinckrodt was filling peculiar (i.e., suspicious) orders telling Mallinckrodt succinctly: "If you think it is suspicious, don't fill it."¹³⁵⁸ Mallinckrodt, however, misconstrued Mr. Kleissie's statement as "if you're going to ship it, it's not suspicious."¹³⁵⁹

Therefore, even though the regulatory requirements and expectations surrounding controlled substances compliance were well established by 2008, Mallinckrodt selectively interpreted those requirements and expectations in such a manner so not to impact their continued distribution of opioid products, although doing so rendered its anti-diversion program ineffective.

In addition to the issues noted above, from the outset, the peculiar order algorithm was highlighting too many orders as peculiar. In April 2009, Ms. Harper wrote, "[w]e have working algorithms, and J. Rausch has been reviewing peculiar orders for several weeks. I have a meeting with Jim tomorrow because **the review is taking several hours a day**, yet still results in him making **a judgment call that he is not comfortable with**."¹³⁶⁰ Later in July 2009, the SOM Team Activity Log noted that the "[p]eculiar order report being generated based upon algorithm settings is 49 pages long."¹³⁶¹

¹³⁵³ Email from Karen Harper to James Rausch *et al.* Re Oxycodone HCL (Jul. 13, 2010), MNK-T1_0000265505.

¹³⁵⁴ See Letter from DEA Deputy Asst. Administrator Joseph Rannazzisi to Mallinckrodt, (Dec. 27, 2007), MNK-T1_0007146632.

¹³⁵⁵ See J. Rausch Deposition at 194:3-23; K. Harper Deposition at 203:1-17.

¹³⁵⁶ See SOM Team Activity Log at 3, MNK-T1_0000477889 (quoting an email from Cathy Stewart in July 2009 "[w]e need to investigate and make sure they're not just gradually increasing their order quantities to not get caught by the [REDACTED] formula threshold.").

¹³⁵⁷ See J. Rannazzisi Letter to All Registrants at 2 (Jul. 27, 2007), MNK-T1_0007146632 ("[R]egistrants that routinely report suspicious orders yet fill these orders without first determining that order is not being diverted into other than legitimate medical, scientific, and industrial channels, may be failing to maintain effective controls against diversion.")

¹³⁵⁸ See Email from B. Ratliff to J. Rausch (Apr. 1, 2008) (K. Harper was copied on that email and Mr. Kleissie was quoted in it.), MNK-T1_0000268860.

¹³⁵⁹ See K. Harper Deposition at 258:14-259:3.

¹³⁶⁰ See Email from Karen Harper to Eileen Spaulding Re Suspicious order monitoring (Apr. 29, 2010) (emphasis added), MNK-T1_0000279142.

¹³⁶¹ See SOM Team Activity Log at 4, MNK-T1_0000477889.

As a result of this ongoing “administrative burden” from having too many orders to review,¹³⁶² Mallinckrodt resorted to simply turning off a portion of the algorithm sometime in 2009 through April 2010.¹³⁶³ Without a fully functioning algorithm, Mallinckrodt had fewer orders to review, but this reduced burden came at the cost of increasing the risk that suspicious orders could slip through. In addition, from 2008 to 2009, Mallinckrodt simply disabled its algorithm altogether, thereby limiting the customer services representatives review to determine if an order was peculiar to:

- Verifying that the account is in good standing by referencing the “Do Not Ship List.”
- Assuring that the order is valid.
- Checking the customer DEA registration status
- Verifying customer has provided a DEA 222 form.¹³⁶⁴

If there were any exceptions in the information, the order was deemed “peculiar.”¹³⁶⁵ However, Mallinckrodt knew that confirmation of the customer’s current DEA registration and status and receipt of a DEA 222 form alone were not appropriate in determining whether an order is suspicious.¹³⁶⁶

At the end of April 2010, Mallinckrodt resumed using the full peculiar order algorithm, but had increased the [REDACTED] to reduce the number of peculiar orders identified.¹³⁶⁷ Mallinckrodt implemented this change despite recognizing the already heightened risk that customers could evade detection of their suspicious orders by gradually increasing their order volume to avoid triggering the threshold.¹³⁶⁸ However, rather than increasing the CSC group’s resources and possibly disrupting the flow of sales and revenue, Mallinckrodt deemed the risk acceptable and memorialized the change in the October 2010 version of the SOM procedure.¹³⁶⁹

¹³⁶² See K. Harper Deposition at 321:20-25.

¹³⁶³ See SOM Team Activity Log at 4, MNK-T1_0000477889. (“30 Day Cumulative Algorithm is turned off because that specific trigger was not spelled out in HDMA guidance and is inflating the peculiar order count”).

¹³⁶⁴ See Mallinckrodt DEA Compliance Procedure, *Identification and Review of Peculiar Orders, Controlled Substance Suspicious Order Monitoring Program*, Draft 4 (Jul. 15, 2008), MNK-T10000263965.

¹³⁶⁵ *Id.*

¹³⁶⁶ Email from Karen Harper to John Adams *et al.* Re Suspicious Order Monitoring Training Notes from Bulk Narcotic Sales Meeting Presentation (Jun. 6, 2008), MNK-T1_0000419956; see also J. Rausch Deposition at 139:14-140:9 (other than verification of the 222 forms Mallinckrodt did not have a suspicious order program in place between fall of 2008 and 2009); William Ratliff Deposition, 242:2-9 (Dec. 19, 2018) (agreeing that it would not be proper to just rely solely on registration status and the 222 form in determining whether or not to ship an order).

¹³⁶⁷ See K. Harper Deposition 321:11-25 (justifying moving from a [REDACTED] formula because the peculiar order report was “too lengthy” and was creating too much of an “administrative burden”).

¹³⁶⁸ See SOM Team Activity Log, MNK-T1_0000477889 (providing a timeline of the SOM program and noting concern with customers gradually increasing their orders).

¹³⁶⁹ See Global Controlled Substance Compliance Procedure, *Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program, C/S Comp. 3.0* (Oct. 29, 2010), MNK-T1_0000264260.

On November 1, 2012, Mallinckrodt changed its procedures for identifying suspicious orders again. Mallinckrodt's new system created separate controls for Oxy 15 and 30 sales to large distributors ("Tier 1") and changed to [REDACTED] its "standard" algorithm for other sales ("Tier 2"):

Tier 1: A monthly limit set annually for large volume wholesalers' or distributors' orders of Oxycodone 15 mg and 30mg SKUs. These products have been identified by DEA as particularly subject to diversion and abuse.

Tier 2: A standard algorithm with respect to volume which sets a

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B. Howard Davis' Concerns

In late 2010, Mallinckrodt retained a former DEA employee, Howard Davis, to specifically evaluate its SOM program.¹³⁷¹ As part of his work assisting Mallinckrodt in evaluating its SOM program, Mr. Davis authored a memorandum in November 2010 that was critical of Mallinckrodt's then current SOM approach.¹³⁷²

Mr. Davis highlighted his concerns about Mallinckrodt’s algorithm, and his feeling that when Mallinckrodt applied the formula in certain situations, it “would be unnecessarily exposing itself to potential liability.”¹³⁷³ Specifically, Mr. Davis was concerned that Mallinckrodt was relying too heavily on the numeric formula while not investing a corresponding amount of effort into investigating orders exceeding the threshold levels (a.k.a. order due diligence).¹³⁷⁴ As Mr. Davis outlined for Mallinckrodt:

Numeric formulas do not identify circumstances that might be indicative of diversion such as: ordering larger quantities of a limited variety regularly that would otherwise not be viewed as suspicious (like ordering controlled substances with few, if any, other drugs or products whether controlled or non-controlled); ordering highly abused controlled substances in limited quantities disproportionate to other products; or even ordering the same controlled substances from multiple suppliers.¹³⁷⁵

To further support his conclusion, Mr. Davis highlighted previous DEA Guidance concerning a manufacturer's "Know Your Customer" obligations as well as recent enforcement actions.¹³⁷⁶

¹³⁷⁰ See Covidien Procedure, *HZQS – Identification, Investigation, and Reports of Controlled Substances Suspicious Orders*, 1 (Nov. 1, 2012), MNK-T1 0005620500.

¹³⁷¹ See Consulting Agreement between Mallinckrodt and Howard Davis (Oct. 18, 2010), MNK-T1_0000427406.

¹³⁷² Memorandum from Howard Davis to Karen Harper, *Suspicious Order Monitoring Program No. C/S Comp 3.0*, (Nov. 2, 2010), MNK-T1 0000269399.

¹³⁷³ *Id.* at 1.

¹³⁷⁴ See *id.* (“DEA has determined that algorithms and/or arithmetic formulas to define whether an order is suspicious may be failing to detect suspicious orders”).

¹³⁷⁵ *Id.* at 2.

¹³⁷⁶ See *id.* at 1; see generally DEA Guidance discussion *infra* at Section 5.3.

In order to remedy the observed deficiencies, Mr. Davis recommended that Mallinckrodt immediately revise its Global Controlled Substance Procedure entitled “Identification and Review of Peculiar Orders” to include criteria “such that a more vigilant determination can be made whether the order is suspicious and/or excessive prior to filling any order, in concert with numeric formulas already in place as the Company deems appropriate.”¹³⁷⁷ By doing so, Mallinckrodt would evaluate the “totality of the circumstances when evaluating an order prior to it being filled, just as the DEA will do in determining if the registrant is operating within the public interest”¹³⁷⁸

Mr. Davis recommended adding the following customer inquiries to Mallinckrodt’s program:

- What percentage of business do controlled substances constitute?
- Is the customer complying with other federal, State, local requirements/laws?
- Is the customer soliciting and/or supplying customers via the Internet unlawfully?
- Does the customer have a business relationship with a physician or other business partner who authorizes prescriptions over the Internet?
- Are prescribing practitioners licenses in the State where the controlled substances are being shipped?
- Are shipments made where the majority of the ultimate customers are serviced from the same practitioner who routinely authorizes “cocktail” prescriptions, i.e., drug combinations that compound or have an antagonistic effect?
- Are controlled substances sold to retail outlets that further dispense them without a prescription?
- Does the customer sell controlled substances for market prices of well above state-of-the-art pricing modules?
- Are sells [sic] made to customers, or their customers, on a cash only basis or are insurance plans accepted?¹³⁷⁹

He also went further and provided Mallinckrodt with a roadmap in the form of a draft SOP.¹³⁸⁰

After receiving the report, Ms. Harper directed Mr. Davis to “revise the QSP Order Monitoring document, incorporate recent program enhancements with recent activities.”¹³⁸¹ Mr. Davis complied and provided some modifications to the procedure that including adding to the “Overview” section the following statements that the procedure was used to:

¹³⁷⁷ *Id.* at 2.

¹³⁷⁸ *Id.*

¹³⁷⁹ *Id.*

¹³⁸⁰ See *Standard Operating Procedure, Due diligence procedures and monitoring of controlled substances sales*, (undated), MNK-Ti_0000269401.

¹³⁸¹ See Email from K. Harper to H. Davis Re Revision of QSP Order Monitoring Attached, (Nov. 18, 2010), MNK-T1_0000264240 (referring to the Global Controlled Substances Compliance Procedure entitled “Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program,” C/S Comp. 3.0, MNK-T1_0000264260).

- Insure (sic) that the chargeback system exists to reimburse wholesalers for discounted pricing to collective buying groups so it does not otherwise flag an order as suspicious
- Insure (sic) that the customer ultimately receiving controlled substances through a buying group has supplied documentation validating any order placed prior to shipment.¹³⁸²

Within approximately two weeks of receiving Howard Davis' critical memo, it appears Karen Harper and Mallinckrodt were preparing to fire Mr. Davis. In an internal email dated November 17, 2010, Ms. Harper characterized Mr. Davis's termination as "Mallinckrodt 'firing' Howard 'quitting'" and she drafted talking points to characterize his departure in terms such as "we are going in a different direction with the program" and "have to do more with less resources."¹³⁸³ In January 2011, Ms. Harper told Mr. Davis that Mallinckrodt would not need his "services in the near future," and she disconnected his laptop and telephone.¹³⁸⁴

In the weeks and months after Mr. Davis presented his memorandum to Mallinckrodt, the CSC group issued a multitude of new procedures in various drafts and versions, including:

- C/S Comp. 2.0 - New Customer Account Set-up, Existing Account & Ongoing Review¹³⁸⁵
- C/S Comp. 4.0 - Customer Audit Program¹³⁸⁶
- C/S Comp. 5.0 - Identification & Review of Suspicious Customer Accts.¹³⁸⁷

14.5.4 Mallinckrodt failed to effectively know their customers' customer despite having enough information to create a robust system to do so.

Mallinckrodt understood the importance of knowing your customer's customer as far back as 2008 and were informed that the DEA expected this of Mallinckrodt in the summer of 2010. DEA, during a July inspection to the Hobart facility, requested an "impromptu meeting" with Mallinckrodt and Ms. Harper.¹³⁸⁸ At the meeting, the DEA Diversion Group Supervisor informed Ms. Harper that the DEA was pursuing a "new direction

¹³⁸² See Global Controlled Substances Compliance Procedure, *Identification and Review of Peculiar Orders Controlled Substance Suspicious Order Monitoring Program*, C/S Comp. 3.0 (undated with handwritten notation "Howard Davis Revision"), MNK-T1_0000269954 ["Davis Revision to C/S Comp. 3.0"]

¹³⁸³ Internal notes prepared by Karen Harper (MNK-T1_0000280821).

¹³⁸⁴ See Email from K. Harper to D. Hunter, Consultant (Howard Davis) Desk Phone and Laptop on Z-4, (Jan. 18, 2011), MNK-T1_0000281479. However, as Mallinckrodt was firing Mr. Davis, the market for Oxycodone 15 and 30 mg appeared to be at or near its peak. See *Oxy Florida Sales*, MNK-T1_0000289708 (Sept. 14, 2010); See Presentation by Mallinckrodt, *Generics Demand Review* Presentation, slide 34 (Jul. 7, 2010), MNK-T1_0007097723.

¹³⁸⁵ See Global Controlled Substances Compliance Procedure, *New Customer Account Set-up and Existing Customer Account Ongoing Review*, (Jan. 4, 2011), MNK-T1_0000264279; MNK-T1_0000264209 (Mar. 28, 2011), MNK-T1_0000259157 (Aug. 8, 2011).

¹³⁸⁶ See Global Controlled Substance Compliance Procedure, *Customer Audit Program*, (Jan. 4, 2011), MNK-T1_0000264214; MNK-T1_0000264205 (Mar. 28, 2011), MNK-T1_0000259162 (Aug. 8, 2011).

¹³⁸⁷ See Covidien, *Identification and Review of Suspicious Customer Accounts*, (Aug. 8, 2011); MNK-T1_0000259153.

¹³⁸⁸ See Email from K. Harper to T. Berry, Re DEA Dialogue 07/20/10 – Suspicious Order Monitoring and Harvard Drug License Suspension, (Jul. 21, 2010), MNK-T1_000269747.

initiative’ whereby enforcement action will be aimed at all entities within the supply chain, including manufacturing registrants.”¹³⁸⁹

Therefore, as Ms. Harper learned from the DEA “[t]he expectation is becoming that suppliers have not only an obligation to know their customers but an additional responsibility to know their customers.” According to Ms. Harper, this was particularly important to the DEA because actions taken against Florida pain clinics for oxycodone diversion were causing those clinics to relocate to Georgia, Ohio, and Texas.¹³⁹⁰

A. The SOM Customer Checklist (a.k.a. SOM Customer Questionnaire)

Another important control that Mallinckrodt employed for anti-diversion purposes was the Suspicious Order Monitoring Customer Checklist. This document was provided to new accounts when they were first set-up and annually to all existing customers.¹³⁹¹ This checklist constituted one of Mallinckrodt’s primary sources of information about its customers and was deployed in July 2009.¹³⁹²

Steven Becker, one of the NAMs, was asked to help put together the questionnaire.¹³⁹³ The intent behind it was to provide it to Mallinckrodt’s customers and confirm that they were looking at various factors that would help them “identify suspicious orders.”¹³⁹⁴ The factors in the questionnaire Becker helped prepare – for example the ordering of excessive quantities of controlled substances and the ordering of controlled substances in quantities disproportionate to non-controlled medications – were based on the Rannazzisi letters, and by Becker’s own admission applied to all of his customers.¹³⁹⁵

After the July 2010 “impromptu meeting,” Ms. Harper and CSC Group changed the SOM Customer Questionnaire to remove the question on whether “our customers monitor their customers.”¹³⁹⁶ In an August 2010 email communicating the change, Ms. Harper explained that the SOM Team’s decision was based on the fact that “there is no actual regulatory obligation to monitor customers’ customer.”¹³⁹⁷ In an ironic juxtaposition of roles, Kate Muhlenkamp, Covidien Product Manager, advocated for retaining the question noting that “[f]rom our perspective (Sales & Marketing), we think it would be a very good idea to require it

¹³⁸⁹ *Id.*

¹³⁹⁰ *Id.*

¹³⁹¹ See Global Controlled Substance Compliance Procedure, *New Customer Account Set-up and Existing Customer Account Ongoing Review*, C/S Comp. 2.0, 2 (Jan 4, 2011), MNK-T1_0000264280.

¹³⁹² See SOM Team Activity Log at 2, MKN-T1_0000477889 (“072009 CDIG begins sending out SOM customer questionnaires.”).

¹³⁹³ S. Becker Deposition at 349:21-24.

¹³⁹⁴ S. Becker Deposition at 351:4-11.

¹³⁹⁵ S. Becker Deposition at 369:9-370:15, 371:14-372:18.

¹³⁹⁶ See Email from K. Harper to G. Collier and K. Muhlenkamp Re Distributors Protocol for Vetting Customers (Aug. 26, 2010), MNK-T1_0000368388.

¹³⁹⁷ *Id.*

despite it not being a regulatory obligation.”¹³⁹⁸ However, Ms. Harper did not agree nor did she see the removal of the question as a problem although it effectively diluted the effectiveness of the questionnaire.¹³⁹⁹

Ms. Harper removed this question despite the fact she was aware that DEA expected Mallinckrodt to undertake KYCC.¹⁴⁰⁰ Given the fact both the DEA and Mallinckrodt’s own sales and marketing team thought knowing whether Mallinckrodt’s customers monitored their customers was important, removing the question from the SOM Customer Checklist was the act of an imprudent company which did not view controlled substances compliance as important. Furthermore, it is troubling when the appointed gatekeepers (compliance) endeavor to dilute the very controls they are charged with protecting.

However, in the end, removal of this question potentially had little impact because Mallinckrodt’s Customer Data Integrity Group (“CDIG”), which had responsibility for ensuring that Mallinckrodt’s customers completed the questionnaire, simply allowed Customer Service to renew accounts even though they failed to complete the form. This prompted Ms. Harper in February 2011 to write Eileen Spaulding, a Compliance Analyst in the Mallinckrodt Dosage Group:

I think I discovered a disconnect in the system. We have significant gaps in that, although CDIG send out the Annual Update SOM Customer Checklist when the system indicates customer account DEA registration is nearing renewal time, THEY DO NOTHING IF THE SOM CUSTOMER CHECKLIST IS NOT EVER RETURNED BY THE CUSTOMER, CDIG primary concern is the license renewal. ... sometimes CSR[s] [Customer Service Representatives] update the license renewal info ... and then CDIG NEVER SENDS AN ANNUAL CHECKLIST.¹⁴⁰¹

Mallinckrodt was aware that DEA expected all registrants to know their customers and in the case of manufacturers that companies were expected to know their customers’ customer. This was a fundamental part of a reasonable and effective SOM program. Both Ms. Harper and Mr. Ratliff, former Chief Security Officer, acknowledged that knowing your customer’s customer is a component of an effective SOM program,¹⁴⁰² and Mr. Ratliff conceded that as early as June 2008 Mallinckrodt understood the importance of knowing your customers’ customers.¹⁴⁰³ Yet despite this knowledge and the explicit feedback to the company by the DEA, Mallinckrodt failed to incorporate a credible KYCC framework into its anti-diversion program.

¹³⁹⁸ See Email from K. Muhlenkamp to K. Harper and G. Collier Re Distributors Protocol for Vetting Customers (Aug. 26, 2010), MNK-T1_0000368390.

¹³⁹⁹ See K. Harper Deposition at 344:22-345:6 (stating that removing the question “was not a detriment to the program”).

¹⁴⁰⁰ *Id.* at 345:7-20.

¹⁴⁰¹ Email from K. Harper to E. Spaulding Re DEA Registration verification, (Feb. 11, 2011), MNK-T1_0000372333 (emphasis in original).

¹⁴⁰² See K. Harper Deposition at 79:10-80:5; W. Ratliff Deposition at 107:4-16.

¹⁴⁰³ See W. Ratliff Deposition at 104:14-106:2.

B. Chargeback Data

Chargebacks are a common pharmaceutical tool used by manufacturers to make distributors “whole” when they sell pharmaceuticals to pharmacies at prices below what the distributor paid to the manufacturer.¹⁴⁰⁴ Since chargebacks represent a loss to the pharmaceutical manufacturer, the industry expends significant resources on its systems and processes to minimize chargebacks and ensure the legitimacy of chargeback claims.¹⁴⁰⁵ This was true in Mallinckrodt’s situation.

In order to interrogate the legitimacy of a chargeback claim, the manufacturer requires specific data on the transaction including the actual customers and the discounted levels at which products were sold to pharmacy customers by the company’s distributors to their customers. As a result, since at least 1998, Mallinckrodt maintained an extensive chargeback database that traced with great granularity sales to the customers of Mallinckrodt’s distributors.¹⁴⁰⁶

According to Ms. Harper:

Mallinckrodt sells controlled substances to wholesalers at a standard price. Some pharmacies negotiate a discounted price. When the wholesaler honors the discounted price to the pharmacy, they then submit a charge back request retroactively to Mallinckrodt so that they can be made financially whole for the difference in price. In doing so the **wholesaler tells Mallinckrodt exactly which pharmacy to which the drugs were sold, what the DEA registration number is, the pharmacy address, the quantity, and which drugs they have sold to that pharmacy.**¹⁴⁰⁷

With access to the chargeback data, Mallinckrodt personnel, including the NAMs, who functioned as the “boots on the ground” for the compliance department, had the downstream visibility that the chargeback data provided to both direct and indirect customers.¹⁴⁰⁸ Mallinckrodt had chargeback data for virtually all transactions with its wholesale distributor customers.¹⁴⁰⁹ This data essentially gave Mallinckrodt visibility into entire supply chain for its products, from manufacturer to distributor to pharmacies and other end dispensers.

¹⁴⁰⁴ See Tyler Lacoma, *What is a Pharmaceutical Chargeback?*, BIZFLUENT (updated Nov. 21, 2018), <https://bizfluent.com/info-8783464-pharmaceutical-chargeback.html>.

¹⁴⁰⁵ *Id.*

¹⁴⁰⁶ See MNK-T1_0007965587-7965588 (chargeback data for 1998-2018 produced by Mallinckrodt in this litigation).

¹⁴⁰⁷ See Deposition of Karen Harper at 11:15-25 (Nov. 7, 2013), Harper Exhibit 9; Apothecary Corporation d/b/a Island Drug v. City of Marco Island, Florida, Case No. 2:10CV-392FtM-36-DNF (M.D. Fla. 2013).) K. Harper Deposition at 227:13-18. In correspondence with the DEA, Ms. Harper described how Mallinckrodt obtains chargeback data and that “distributors must provide Mallinckrodt with specific detailed information indicating how much product was sold to each end user pharmacy.”; *see also*, Letter from K. Harper Letter to P. Kleissle (Nov. 1, 2010), MNK-T1_0000280607.

¹⁴⁰⁸ See, e.g., V. Borelli Deposition at 130:5-136:24; S. Becker Deposition at 158:13-18, 161:24-163:1.

¹⁴⁰⁹ See K. Harper Deposition at 229:15-18. Ms. Harper informed the DEA that Mallinckrodt “assumes most transactions would result in a chargeback request” by a distributor. See Email from K. Harper to K. Hamilton Re Explanation of Mallinckrodt Chargeback System (Nov. 2, 2010), MNK-T1_0000387492. For example, 96% of all Oxy 15 orders and 98% of all Oxy 30 orders were subject to chargeback requests, and hence would be in the chargeback database. J. Gillies (30)(b)(6) Deposition at 271:3-274:2.

While Mallinckrodt was aware of the power of chargeback data in 2007, it was not until the 2009-2010 timeframe that the CSC Group became interested in using the data.¹⁴¹⁰ Even then, Ms. Harper was slow to utilize chargeback data – in fact, despite acknowledging that it would not be complicated to pull the chargeback data and receive training for how to utilize it, it took Ms. Harper at least four months between when she first asked about the chargeback data in late 2009 and when she actually purportedly took action on it later in 2010.¹⁴¹¹

In another ironic juxtaposition of roles, Ginger Collier, the Director of Marketing for generics, related that she directed Kate Muhlenkamp Neely, the product manager for oxycodone, to analyze chargeback data relating to the sale of opioid products to doctors in Florida.¹⁴¹² The data revealed that many pharmacies and pain clinics, particularly in Florida, were buying from multiple distributors (which is a red flag for diversion). Ms. Collier found this surprising based on her experience in the pharmaceuticals industry.¹⁴¹³

When Mallinckrodt subsequently notified distributors that it would not pay chargebacks on sales to multi-distributor customers, Mallinckrodt failed to report any of the orders that gave rise to multi-distributor sales to the DEA as suspicious.¹⁴¹⁴ Simply put, once Mallinckrodt identified pharmacies that were acquiring opioid products from multiple distributors, it moved quickly to refuse to pay chargebacks on those sales (thereby saving the company money), but failed to report these orders as suspicious to the DEA. Furthermore, by not employing the chargeback data the company had access to in a timely fashion, Mallinckrodt missed an opportunity to significantly improve its suspicious order monitoring program.

C. Chargebacks & DEA Enforcement Actions

During the review period, the DEA investigated and shut down several Mallinckrodt wholesalers including Harvard Drug Group; Masters Pharmaceuticals, Inc.; Keysource Medical; Kinray, LLC.; Value Drug, Inc.; and Sunrise Wholesalers, Inc. When taken together, the following pattern emerged:

1. These were not the Big Three distributors.
2. All were distributing high volumes of opioid products (e.g., Keysource Medical distributing 48 million dosage units to Florida customers).
3. All were distributing to multiple pharmacies (e.g., Kinray distributing to more than 20 New York pharmacies) which were filling opioid prescriptions for other than legitimate medical purposes; and

¹⁴¹⁰ See Email from Karen Harper to herself (Nov. 18, 2010), MNK-T1_0000280835 (Harper notes indicating that she was currently working on adding chargeback data analysis to the SOM program.); Harper deposition at 355:22-356:22.

¹⁴¹¹ See K. Harper Deposition at 358:21-367:25; see also K. Harper Exhibit 22 (Mar. 29, 2010 email from Karen Harper to Carrie Johnson re chargeback information requests, MNK-T1_0000500657).

¹⁴¹² See G. Collier Deposition at 50:8-24 (Jan. 8, 2019).

¹⁴¹³ See *id.* at 197:6-11 & Collier Ex. 15 (Nov. 11, 2010 email between G. Collier, S. Becker, and V. Borelli, MNK-T1_000418885) & Collier Ex. 16 (Sept. 2010) (Summary of customers sourcing more than 2 distributors for Oxy 30, MNK-T1_000418885).

¹⁴¹⁴ G. Collier Deposition at 236:13-17.

4. All failed to report large numbers of suspicious orders to the DEA and failed maintain effective controls against the diversion of controlled substances.¹⁴¹⁵

In the case of each of these distributors that the DEA investigated and shut down, Mallinckrodt had similar data to the DEA, and where Mallinckrodt was the sole supplier of opioid products, the company had access to the same information as the DEA demonstrating significant diversionary activities by these wholesalers.

Mr. Becker, one of Mallinckrodt's NAMs, who had responsibility for many of the customers targeted by the DEA, related that:

- (a) in the case of the Harvard Drug Group, he had the same information as the DEA regarding Mallinckrodt sales and could have conducted his own analysis on them, but did not because he was not suspicious of Harvard until he saw the press release indicating that its license was suspended;¹⁴¹⁶
- (b) that he would have found it "alarming" that Mallinckrodt's chargeback data showed that Harvard, doing business as a vet supply company (First Veterinary Supply), supplied 12,487 orders of Oxy 15 and Oxy 30 to doctors, 92.4% of which went to the state of Florida, but that he was unaware of this information even though it was clear from the chargeback data;¹⁴¹⁷
- (c) for Value Drug, he had access to the chargeback data showing Value Drug's sales to downstream customers, but never requested it from headquarters to evaluate them;¹⁴¹⁸
- (d) with respect to Masters Pharmaceuticals, he had access to information regarding Masters' sales in the chargeback system, but did not recall having any suspicions about Masters until the DEA suspended its license;¹⁴¹⁹
- (e) in the case of Kinray, he also had access to detailed information regarding Kinray sales and while he evaluated them, Kinray's activities did not raise any concerns for him.;¹⁴²⁰ and
- (f) finally, with respect to Keysource, he conceded that the product monitoring group should have evaluated sales data for Keysource, but that he does not know if they ever did.¹⁴²¹

Mallinckrodt therefore had access to much, if not all, of the information possessed by the DEA, but Mallinckrodt failed to make use of that data.

¹⁴¹⁵ See S. Becker Deposition Ex. 33 (June 15, 2010) (DEA Press Release re Harvard's suspension); S. Becker Deposition Ex. 36 (Sept. 15, 2015) (DEA Notice of Decision & Order re Masters); S. Becker Deposition Ex. 38 (Jun. 10, 2011) (Press Release re KeySource license suspension Becker Ex. 37 (Dec. 23, 2016) (DOJ Press Release re civil penalty levied against Cardinal); S. Becker Deposition Ex. 35 (Jun. 25, 2014) (DOJ Press Release re Value Drug settlement).

¹⁴¹⁶ See S. Becker Deposition at 312:11-321:12.

¹⁴¹⁷ S. Becker Deposition at 190:4-192:19.

¹⁴¹⁸ *Id.* at 322:14-324:18.

¹⁴¹⁹ *Id.* at 325:1-328:16.

¹⁴²⁰ *Id.* at 329:8-333:6.

¹⁴²¹ *Id.* at 335:1-338:17.

14.6 Accountability - Consistent Enforcement

14.6.1 Even in the face of credible evidence, Mallinckrodt frequently failed to hold its distributors accountable for not having adequate SOM Programs

On August 23, 2011, Mallinckrodt met with the DEA in Washington D.C. During that meeting, the DEA told Mallinckrodt that it was under “tremendous pressure from Congress and the White House” to address the Florida oxycodone problem. The DEA also told Mallinckrodt that its distributors were not sufficiently validating the legitimacy of orders received from pharmacies in Florida.¹⁴²² Therefore, a “disproportionate amount” of the company’s 15 and 30 mg. oxycodone tablets were ending up in Florida pharmacies, and the DEA concluded that Mallinckrodt’s distributors potentially had SOM programs that were inadequate.¹⁴²³

After learning this from the DEA, Mallinckrodt met with Cardinal in September 2011 to discuss the potential problem pharmacies that had purchased large amounts of oxycodone 15 and 30 milligram dosages from Cardinal. Before the meetings with the DEA and Cardinal, Mallinckrodt had generated two “Top 20” lists of such pharmacies – the Top 20 from Florida and the Top 20 in the other states.¹⁴²⁴

After Mallinckrodt and Cardinal met on October 2, 2011, Ms. Harper sent Cardinal revised Top 20 lists that apparently reflected the outcome of the joint review by and negotiations between Mallinckrodt and Cardinal.¹⁴²⁵ The net result was that after discussing these pharmacy situations with Cardinal, Mallinckrodt decided that 8 of the 20 Florida pharmacies should be removed from the chargeback restriction list. For the pharmacies outside of Florida, Mallinckrodt pared the list down to only 6 of the original 20 pharmacies. For the other 14, the notes indicated that the “Cardinal SOM file [was] not yet reviewed by Mallinckrodt,” which suggests Cardinal had provided Mallinckrodt with information and the company was going to permit chargebacks for those pharmacies until it could review the file. Mallinckrodt and Cardinal set a target “cut off” date of October 14, 2011 to complete the review of the Top 20 pharmacies.¹⁴²⁶

Mallinckrodt and Cardinal met again on October 13, 2011.¹⁴²⁷ This meeting appears to have resulted in an additional 7 Florida pharmacies being cleared for chargebacks by Mallinckrodt.¹⁴²⁸ This raised the total of cleared Florida pharmacies to 15 of the original Top 20 (75%) and 11 of 20 (55%) for the non-Florida pharmacies. In sum, Mallinckrodt’s negotiations with Cardinal significantly eroded the effectiveness of

¹⁴²² See Draft Notes for SOM Steering Committee Meeting (Sept. 28, 2011), MNK-T1_0002077756.

¹⁴²³ *Id.*

¹⁴²⁴ See Top 20 spreadsheet (Sept. 21, 2011), MNK-T1_000473333).

¹⁴²⁵ See Email from K. Harper to Michael Moné, *et al.*, Re Revised Chargeback Restriction Listing for Pharmacies Purchasing Mallinckrodt Product from Cardinal, (Oct. 2, 2011), MNKT1_0000472004 (copying Mallinckrodt attorney Donald Lohman and others at Mallinckrodt).

¹⁴²⁶ See Email from Jane Williams to Scott Decker *et al.*, Re Revised Chargeback Restriction Listing for Pharmacies Purchasing Mallinckrodt Product from Cardinal, (Oct. 5, 2011), MNK-T1_0000471988.

¹⁴²⁷ See Email from Donald Lohman to Nicholas Rausch Re Chargeback Status, (Oct. 17, 2011), MNK-T1_0000471975 (referencing prior week’s meeting and clearing by Mallinckrodt of two pharmacies for chargeback reimbursement).

¹⁴²⁸ See Spreadsheet Cardinal Health, (Oct. 13, 2011), MNK-T1_0000460617.

chargeback restrictions – the primary tool Mallinckrodt had used to ensure its distributors were cracking down on the problem pharmacies.

Mallinckrodt also purportedly “audited” all the previously listed distributors the DEA shut down. The CSP group described these audits as “high level,” which appears to be an attempt to justify the lack of care in conducting them.¹⁴²⁹ Mallinckrodt did not inquire about the distributor’s customers, even when it had chargeback data showing unusually high orders of oxycodone or inquire as to whether the distributors had cut off any pharmacies.¹⁴³⁰

The Masters Pharmaceutical audit is an excellent example of Mallinckrodt's lack of diligence. In the Masters audit, completed in December 2010, Mallinckrodt concluded Masters possessed an “adequate” SOM process.¹⁴³¹ Prior to that audit, Masters had identified a problem pharmacy, Brooks Pharmacy, which Masters had cut off in October 2010.¹⁴³² However, apparently Mallinckrodt and Masters did not bother to discuss problem pharmacies such as Brooks during the audit because Mallinckrodt, through Cardinal, continued to supply Brooks with opioids after auditing Masters.¹⁴³³

Furthermore, while the CSC often generated “indirect match” reports tracing Mallinckrodt opioid products to doctors and pharmacies after they had been shut down by the DEA, Mallinckrodt failed to use its data proactively to stop diversion.¹⁴³⁴ As highlighted by Mr. Becker, Mallinckrodt shipped to problematic distributors up until the time the distributors were shut down by the DEA.¹⁴³⁵ Even Ms. Harper acknowledged that pharmacies that been placed on a distributor’s termination list were problematic, and that shipping orders to these pharmacies after placement on such as list “would not be indicative of an effective SOM program.”¹⁴³⁶

14.6.2 Mallinckrodt failed to hold employees accountable for being non-compliant and out of sync with its values.

This lack of accountability can be seen in several ways. First, Mallinckrodt made no effort to assess the extent to which its opioids sales goals and results were dependent on supplying irresponsible distributors and their remote customers. For example, at its national sales meeting for fiscal year 2011, Mallinckrodt celebrated that

¹⁴²⁹ See Email from K. Harper to B. Ratliff Re Pete Kleissle, Oxy Investigation (July 27, 2009), MNK-T1_0000307203.

¹⁴³⁰ See K. Harper Deposition at 419:23-421:14.

¹⁴³¹ See Masters Audit, *Controlled Substance Compliance/Suspicious Order Monitoring Distributor Customer Audit Checklist* at 5, (Dec. 8, 2010), MNK-T1_000029608.

¹⁴³² See Harper Exhibit 29 (Oct. 20, 2011 email from Wayne Corona to Karen Harper re cut-off pharmacies, MNK-T1_0000311741).

¹⁴³³ See K. Harper Deposition at 417:20-419:12; *see also*, Harper Exhibit 30 (Brooks Pharmacy Monthly Total 15mg & 30mg Oxy “Sales Qty Govt UOM” 2010-2011, MNK-T1_0001519959 & MNK-T1_0001810303).

¹⁴³⁴ See Email chain between Carrie Johnson, Karen Harper and others (Sept. 6, 2011), MNK-T1_0000283884 (demonstrating that Indirect Match Reports were based on information re doctors and pharmacies that had already been shut down); Indirect Customer Match Report (Sept. 6, 2011), MNK-T1_0000283884.

¹⁴³⁵ See S. Becker Deposition Ex. 36 (Sept. 15, 2015 DEA Notice of Decision & Order re Masters); S. Becker Deposition Ex. 33 (June 15, 2010 DEA Press Release re Harvard’s suspension).

¹⁴³⁶ See K. Harper Deposition at 416:24-417:16.

generics sales were 99% of its plan and that everyone in the sales forces was “in the money,” most between 100-115% of goals, but failed to even acknowledge that many of its major customers during 2011 had had their DEA licenses suspended or revoked.¹⁴³⁷ I have seen no evidence in the record that anyone at Mallinckrodt was subjected to disciplinary sanction or suffered any negative consequence for selling to any of the DEA suspended or revoked customers.

Second, the head of Mallinckrodt’s compliance department, Karen Harper, testified that she offered her resignation after Mallinckrodt’s settlement with the DOJ and DEA on the grounds that the deficiencies in Mallinckrodt’s SOM program “happened on my watch,” but her resignation was not accepted.¹⁴³⁸

Third, in the case of Victor Borelli, customer Service Manager Cathy Stewart warned her superiors – Karen Harper and Bill Ratliff -- that the customer service representatives all said that “Borelli will tell them anything they want to hear just so he can get a sale.” Furthermore, Mr. Ratliff testified that despite this warning, and despite not knowing whether Borelli was an “honest” or “reliable” person, no steps were taken to verify the information provided by Borelli.¹⁴³⁹



Dr. Seth B. Whitelaw

President & CEO, Whitelaw Compliance Group, LLC.

¹⁴³⁷ See *Specialty Generics National Sales Meeting* presentation at 5 (Nov. 9, 2011), MNK-T1_0002450579; S. Becker Deposition at 152:1-154:3 (discussion of Exhibit 12).

¹⁴³⁸ See K. Harper Deposition at 98:05-13.

¹⁴³⁹ W. Ratliff Deposition at 284:13-285:10, 291:16-23 (“Q: Did – did anyone do anything to try to verify Mr. Borelli’s information about Sunrise? A: I don’t know. Q: In your opinion, should something have been done to try to verify this information? A: Based on what we know now, yes.”)

Appendices



Appendix A: Opioid Public Health Crisis – A Brief Discussion

Figure 1: Opioid Public Health Crisis – A Brief Discussion

A recent House Energy and Commerce Committee Report declared that the “opioid epidemic is the worst drug crisis in America’s history.”¹⁴⁴⁰ Furthermore, the report noted that “[a]ccording to the Centers for Disease Control and Prevention, more than 351,000 lives have been lost to opioid overdoses since 1999, with no signs of abating.”¹⁴⁴¹ In fact, the report concluded the epidemic has reached a point that it has “helped drive a decline in the U.S. life expectancy at a time when life expectancy is improving in many places around the world.”¹⁴⁴² Moreover, data from the CDC indicates that people who are addicted to opioids are 40 times more likely to subsequently become addicted to heroin.¹⁴⁴³

The United States also is unique for the volume of opioid medicinal products used. A 2017 report by the International Narcotics Control Board noted that “[i]n 2016, the country with the highest consumption of hydrocodone continued to be the United States, with 33.4 tons, equivalent to 99.1 per cent of total global consumption.”¹⁴⁴⁴ In that same report, it was noted that the United States consumed 72.9 per cent of the world’s total of oxycodone for that same time period.¹⁴⁴⁵ However, the opioid crisis is not just about addiction and overdoses. It also impacts the patients who legitimately need access to opioids.

The opioid epidemic also is not a newly recognized public health crisis. As Judge Polster has written in this case, “[i]t is accurate to describe the opioid epidemic as a man-made plague, 20 years in the making.”¹⁴⁴⁶ Also in August 2001, the House Subcommittee on Oversight and Investigations held a public hearing discussing the legitimate and illegitimate uses of oxycontin.¹⁴⁴⁷ Additionally, in December 2003, the United States General Accounting Office (“GAO”) published for Congress on the abuse, use and diversion of OxyContin.¹⁴⁴⁸

¹⁴⁴⁰See U.S. House Energy & Commerce Committee Report, *Red Flags and Warning Signs Ignored: Opioid Distribution and Enforcement Concerns in West Virginia*, 115th Cong., 4 (Dec. 19, 2018).

¹⁴⁴¹ *Id.* (quoting information provided by the Centers for Disease Control, citations omitted).

¹⁴⁴² *Id.*

¹⁴⁴³ See Centers for Disease Control and Prevention, *Vital Signs: Today’s Heroin Epidemic* (Jul. 2015)

¹⁴⁴⁴ See Int’l Narcotics Control Bd., *Narcotic Drugs: Estimated World Requirements for 2018; Statistics for 2016*, . 36 (2017).

¹⁴⁴⁵ *Id.* at 37.

¹⁴⁴⁶ See Amanda Bronstad, Opioid Judge Refuses to Dismiss Claims That Drug Companies Caused ‘Man-Made Plague,’ LAW.COM (Dec. 20, 2018 at 04:35 P.M.), <https://www.law.com/2018/12/20/opioid-judge-refuses-to-dismiss-claims-that-drug-companies-caused-man-made-plague/>.

¹⁴⁴⁷ See *Oxycontin: Its Use and Abuse, Hearing Before the House Subcomm. on Oversight and Investigations, H. Comm. on Energy and Commerce*, 107 Cong., (Aug. 28, 2001), <https://www.govinfo.gov/content/pkg/CHRG-107hhrg75754/html/CHRG-107hhrg75754.htm>

¹⁴⁴⁸ See *Oxycontin: Its Use and Abuse, Hearing Before the House Subcomm. on Oversight and Investigations, H. Comm. on Energy and Commerce*, 107 Cong., (Aug. 28, 2001), <https://www.govinfo.gov/content/pkg/CHRG-107hhrg75754/html/CHRG-107hhrg75754.htm>; see also U.S. Gen’l Accounting Office, *Prescription Drugs: OxyContin Abuse and Diversion and Efforts to Address the Problem*, GAO-04-110 (Dec. 2003), <https://www.gao.gov/new.items/d04110.pdf>.

The State of Ohio also has been particularly hard-hit by this crisis. In 2010, a report by the Ohio Prescription Drug Abuse Task Force noted that “[i]n 2007, unintentional drug overdose surpassed motor vehicle crashes and suicide as the leading cause of injury death in Ohio for the first time on record.”¹⁴⁴⁹ The report further noted that “prescription opioids are largely responsible for this alarming increase in drug overdose death rates.”¹⁴⁵⁰ In 2012, Ohio was noted as being one of the top 10 states for pharmacy dispensing of oxycodone (#5), hydrocodone (#7), hydromorphone (#8), and oxymorphone (#7).¹⁴⁵¹ For oxycodone, Ohio only lagged behind New York, California, Pennsylvania and Florida.¹⁴⁵²

The opioid crisis is also unique in that activities in one location can have an impact on jurisdictions many miles or even states removed. Thus, the migration of opioids from the originating pharmacy to other cities, counties and states in some ways resembles Prohibition era bootlegging. Like alcohol bootlegging before it, opioid diversion takes a risk mitigation approach moving from areas with strong enforcement to areas that are weaker. Consequently, while Florida started out as “ground zero” for diversionary pharmacies, those pharmacies ultimately spread to others states such as Kentucky, West Virginia and Ohio.¹⁴⁵³

With the purchase of the product tied to one location and consumption tied to another, opioid migration also resembles Prohibition era bootlegging. This phenomenon, the so-called “oxy express,” describes the frequent trips made by thousands of individuals to states like Florida to purchase opioids easily and take them back to the State where they reside¹⁴⁵⁴ This phenomenon has been documented and commented on by both the DEA, as well as the distributors and manufacturers of opioid products.¹⁴⁵⁵

By failing to monitor the number of opioid dosage units actually being distributed in a location versus what was legitimately needed, the lack of suspicious order due diligence on the part of opioid product manufacturers and distributors certainly has contributed to the ability of opioid products to move with relative ease between locales.

Therefore, as the objective evidence shows, the wide-spread use together with the accompanying abuse of prescription opioids constitutes an important, chronic public health crisis.

¹⁴⁴⁹ See OHIO PRESCRIPTION DRUG ABUSE TASK FORCE; FINAL REPORT TASK FORCE RECOMMENDATION, 19 (Oct. 1, 2010).

¹⁴⁵⁰ *Id.* at 20.

¹⁴⁵¹ See Presentation by Krista Peck, Regulatory Update, 12 (circulated Jun. 10, 2014), MCKMDL00403517 at 00403529 (citing source as DEA Distributors Conference Oct. 2013).

¹⁴⁵² *Id.*

¹⁴⁵³ See Presentation by McKesson Corporation, *Prescription Drug Abuse - The National Perspective*, 15 (2014), MCKMDL00407451 at 00407465.

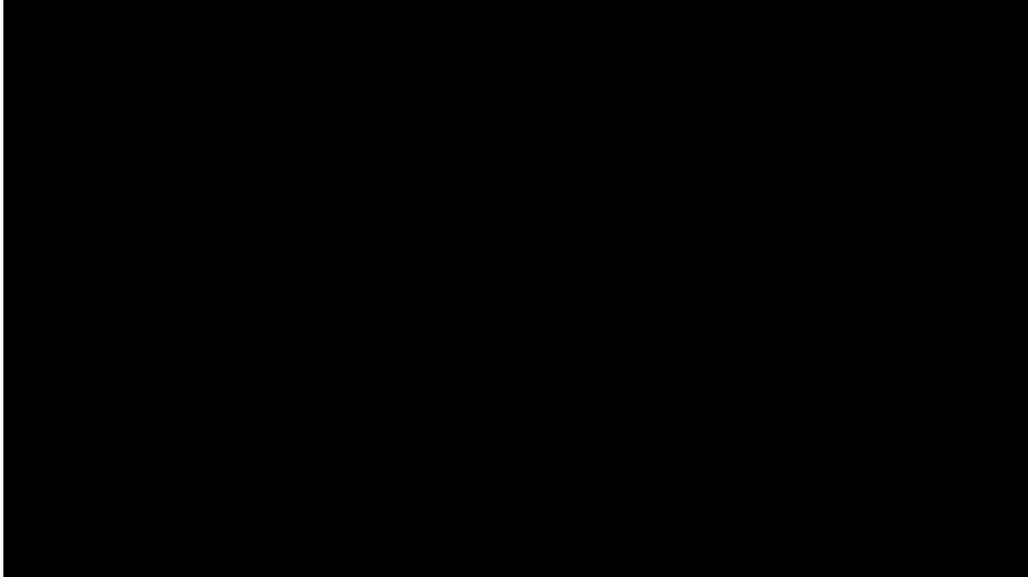
¹⁴⁵⁴ See The ‘Oxy Express’: Florida’s Drug Abuse Epidemic, NPR (Mar. 2, 2011), <https://www.npr.org/2011/03/02/134143813/the-oxy-express-floridas-drug-abuse-epidemic>; Pat Beall, *Florida cuts of oxy: Death, Devastation follow*, THE PALM BEACH POST, <https://heroin.palmbeachpost.com/florida-cuts-off-oxycodone-death-devastation-follow/?ref=lowerTeases>, (last visited Apr. 4, 2019).

¹⁴⁵⁵ See *e.g.*, Karen Harper Depo. at 91:9-92:19; Michael Oriente Depo. at 93:6-94:15.

David Family Pharmacy
705A-809000758
2000 David Avenue
Stark, Copeland County, NY, 44133

CONFIDENTIAL - SUBJECT TO PROTECTIVE ORDER

Oxycodone Distribution to Euclid Family Pharmacy



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March 2019

Figure 2: Oxycodone Distribution to Euclid by Distributor¹⁴⁵⁶

Major Distributors from
January 2005 to January
2018.

¹⁴⁵⁶ See Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated Mar 25, 2019 at 318 (Oxycodone Distribution to Euclid Family Pharmacy).

Appendix B: Applying the Standards

Figure 1: Table of Standard Elements Found in Policies and Procedures¹⁴⁵⁷

<u>General Elements</u>	
<ul style="list-style-type: none"> • Title: identifies the subject • Reference number: useful for internal tracking • Statement of purpose: may provide citations to regulations • Scope: defines resources covered, such as all PHI or all confidential information including proprietary business information • Definitions: defines terms that have special meaning • References: lists any external sources of information or standards • Effective date: the date the policy or procedure was put into place 	<ul style="list-style-type: none"> • Review/revision date: the date of any review and change. Policies should never be destroyed. If a policy is no longer applicable, it should be retired and placed in a permanent file. This is because it may be necessary for the organization or a member of the workforce to demonstrate that a previous action was or was not consistent with the old policy • Authority and approval: identifying who may authorize approval • Rider: may be used to authenticate receipt and agreement to abide by
Policy Statement	<ul style="list-style-type: none"> • Measurable objectives and expectations: these are the primary statements of the policy • Responsibilities: assigns duties for implementation • Compliance enforcement: describes how the policy will be monitored and enforced
Detailed Procedures Steps	<ul style="list-style-type: none"> • Resources: tools and other resources required to perform the procedure • Detailed procedural steps: a list, flowchart, or storyboard outlining the sequence of steps to perform • Associated forms/screens: illustrates data entry or retrieval • Performance expectations: quantity and quality standards

¹⁴⁵⁷ See Margret Amatayakul, *Practical Advice for Effective Policies, Procedures*, 74 J. OF AHIMA.4: 16A-D (Apr. 2003), <http://library.ahima.org/doc?oid=59451#.XBfhIfZFwuW>.

associated with tasks

Figure 2: Controlled Substances Compliance

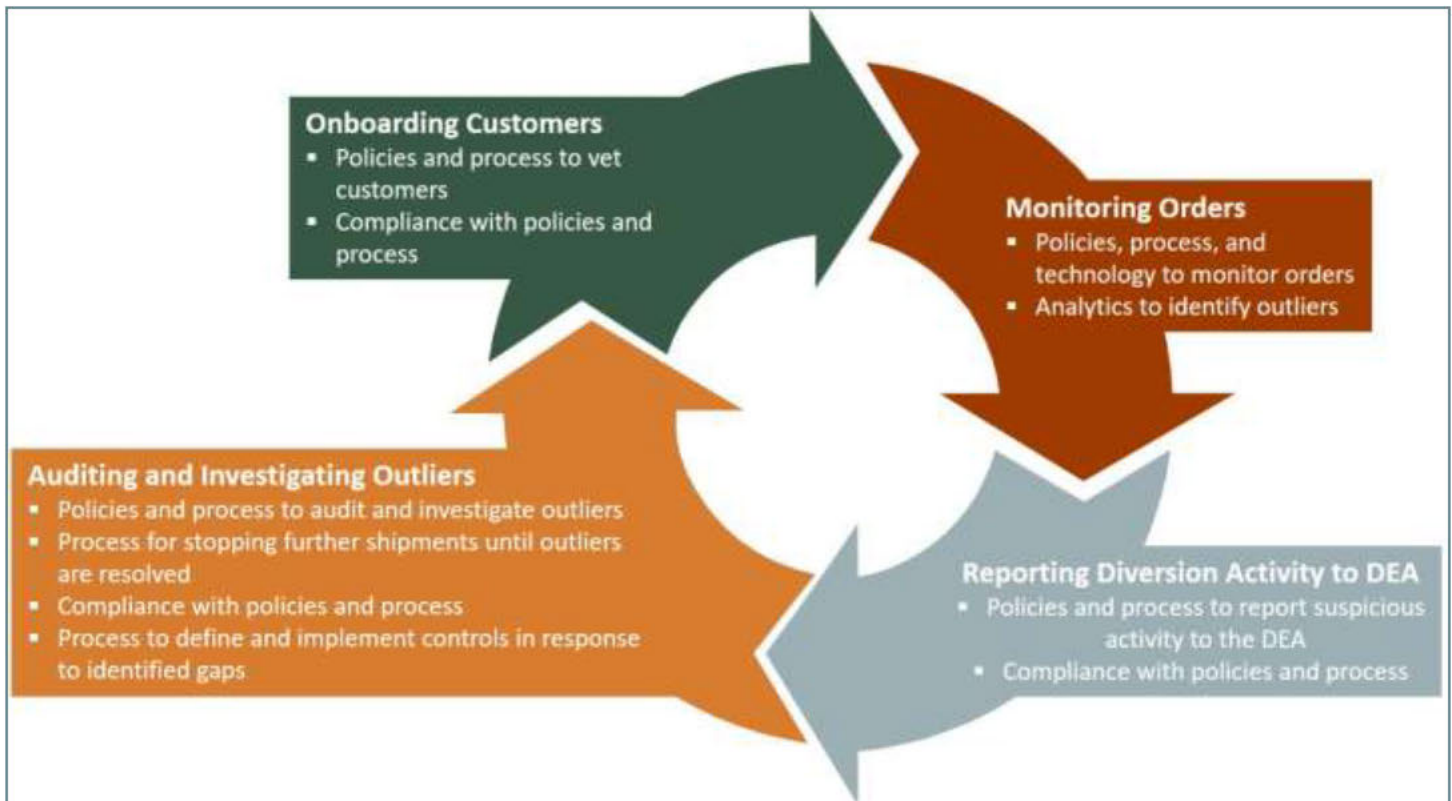


Figure 3: Corporate Compliance



Appendix C: McKesson Key Facts and Figures

Figure 1: McKesson Self-Reported Data Points

DATA Item	YEAR			
	2014	2015	2016	2017
Controlled Substance Program Headcount	30 ¹⁴⁵⁸			44 ¹⁴⁵⁹ (incl. 9 support members)
Number of Distribution Centers	30 ¹⁴⁶⁰			27 ¹⁴⁶¹
Number of Customers (Pharmacies)	~25,000 ¹⁴⁶²	34,816 ¹⁴⁶³		>40,000 ¹⁴⁶⁴
Number of Nightly Line Items Processed from Customers	1.2 million ¹⁴⁶⁵			
Number of Suspicious Orders Reported		230,000 ¹⁴⁶⁶	220,000 ¹⁴⁶⁷	145,000 ¹⁴⁶⁸
Controlled Substances as Percentage of Enterprise-wide Sales		3.1-4.2% ¹⁴⁶⁹	3.1-4.2% ¹⁴⁷⁰	3.1-4.2% ¹⁴⁷¹
Controlled Substances as a Percentage of U.S. Pharm.		4.3-5.2% ¹⁴⁷²	4.3-5.2% ¹⁴⁷³	4.3-5.2% ¹⁴⁷⁴

¹⁴⁵⁸ McKesson Corporation, Presentation to the USAO, Northern D. W. Va. and DEA, 9 (Mar. 12, 2014) MCKMDL00409116 ["USAO Presentation 2014"].

¹⁴⁵⁹ See ISMC Controlled Substances Monitoring Program Operating Manual, Version 1.3., 6, § 3.1, Figure 3, (Jan. 6, 2017), MCKMDL00395206 ["ISMC CSMP Manual 2017"].

¹⁴⁶⁰ See USAO Presentation 2014 at 4.

¹⁴⁶¹ See MCK Teamsters Response at 1.

¹⁴⁶² See USAO Presentation 2014 at 4.

¹⁴⁶³ See Presentation, *McKesson's Controlled Substances Monitoring Program - Regulatory Affairs Training*, 18 (undated), MCKMDL00336532; but see N. Hartle Deposition, 17:20-23 and 18:1-6 (Aug. 1, 2018) (Establishing via metadata that the training deck discussed in the De Gutierrez-Mahoney deposition was produced on December 31, 2015). Customer figure as of September 2015.

¹⁴⁶⁴ See MCK Teamsters Response at 12.

¹⁴⁶⁵ See USAO Presentation 2014 at 4.

¹⁴⁶⁶ See MCK Teamsters Response at 24.

¹⁴⁶⁷ See MCK Teamsters Response at 24.

¹⁴⁶⁸ See MCK Teamsters Response at 24.

¹⁴⁶⁹ See MCK Teamsters Response at 27.

¹⁴⁷⁰ See MCK Teamsters Response at 27.

¹⁴⁷¹ See MCK Teamsters Response at 27.

¹⁴⁷² See MCK Teamsters Response at 27.

¹⁴⁷³ See MCK Teamsters Response at 27.

DATA	YEAR			
Sales				

Headcount	YEAR			
	2014	2015	2016	2017
Controlled Substance Program Headcount	30 ¹⁴⁷⁵			44 ¹⁴⁷⁶ (incl. 9 support members)

Figure 2: McKesson Program Overview (Circa 2015)¹⁴⁷⁷



¹⁴⁷⁴ See MCK Teamsters Response at 27.

¹⁴⁷⁵ McKesson Corporation, Presentation to the USAO, Northern D. W. Va. and DEA, 9 (Mar. 12, 2014) MCKMDL00409116.

¹⁴⁷⁶ See ISMC CSMP Manual 2017 at § 3.1, Figure 3, (Jan. 6, 2017), MCKMDL00395206.

¹⁴⁷⁷ See Presentation, *McKesson's Controlled Substances Monitoring Program - Regulatory Affairs Training*, at 24 (Reproduced by expert from slide).

Appendix D: Cardinal Health Key Facts and Figures

Figure 1: Various Data Points

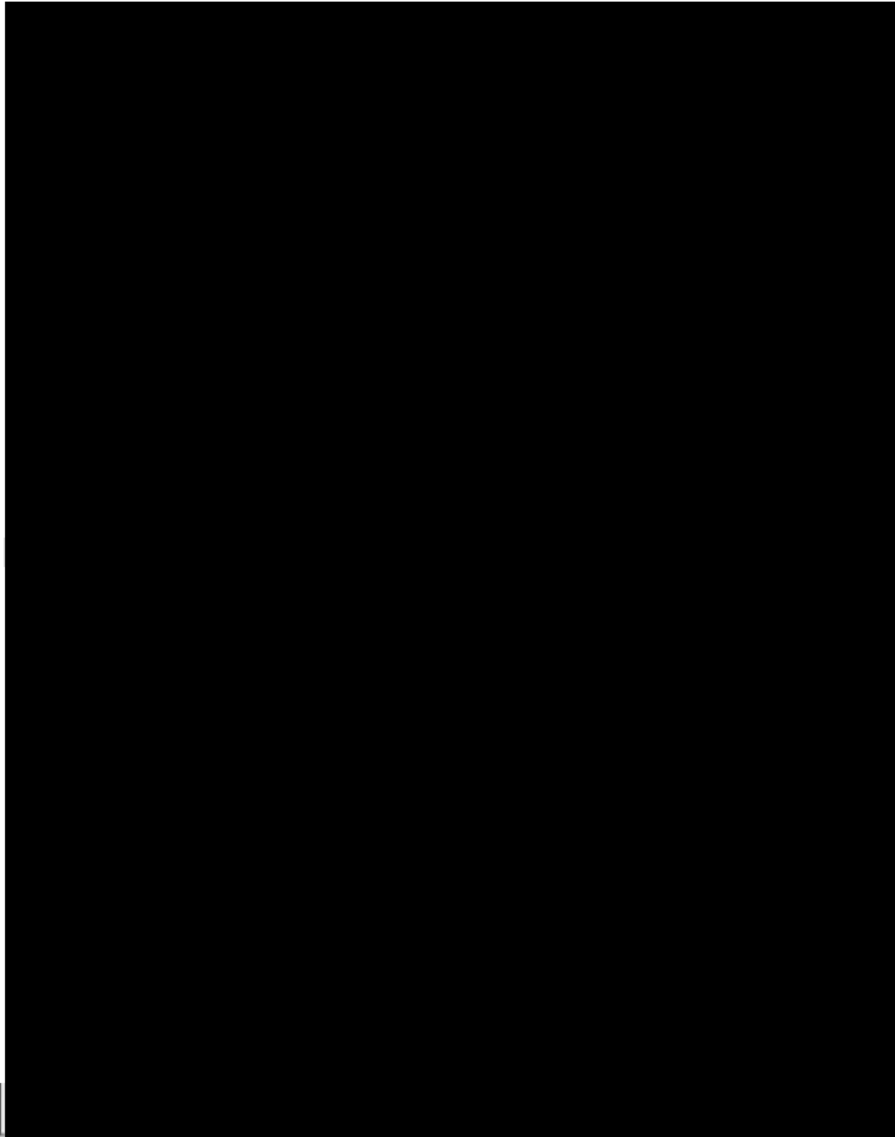
DATA Item	YEAR						
	2007- 2012	2012	2013	2014	2015	2016	2017
Total number of U.S. Customers with shipments suspended or terminated by Cardinal ¹⁴⁷⁸	~330						
W. Va. Suspicious Order Reports Submitted to the DEA ¹⁴⁷⁹	unknown	245	542	557	285	260	181
Dosage Units of Oxycodone & Hydrocodone Shipped to W. Va. (Millions) ¹⁴⁸⁰	174	36	31	32	40	34	--

¹⁴⁷⁸ See W. Va. Red Flags Report at 246.

¹⁴⁷⁹ *Id.* at 243.

¹⁴⁸⁰ *Id.*

Figure 2: Table 5 -Customer Release Percentage¹⁴⁸¹



¹⁴⁸¹ See Cardinal Health, *QRA SOM Customer Analytics General Work Instructions*, 15 (Sept. 20, 2013) (Appendix 5), CAH_MDL2804_00012244 at CAH_MDL2804_00012249, CAH_MDL2804_00012263.

Appendix E: AmerisourceBergen Key Facts and Figures

Figure 1: ABC 2009 Default Thresholds¹⁴⁸²

Customer Type	Definition (Monthly Dollar Volume)	Oxycodone Threshold	Hydrocodone Threshold
Small Retail	Total [REDACTED]	[REDACTED]	[REDACTED]
Medium Retail	Total [REDACTED] and [REDACTED]	[REDACTED]	[REDACTED]
Large Retail	Total [REDACTED]	[REDACTED]	[REDACTED]

Category of Customers: (Average Retail Controlled Substances Ratio [REDACTED])

1. Low dollar volume and a low ratio of controlled substances
2. High dollar volume and a low ratio of controlled substances
3. Low dollar-volume ([REDACTED] – small retail) and high ([REDACTED]) ratio of controlled substances
4. High dollar volume and controlled substances ratio.

Figure 2: ABC Diversion Control Policies and Procedures (SOPs)¹⁴⁸³

Document	Policy or SOP	History & Comments
DCP – 12.1.0 Know Your Customer Due Diligence	Policy	Original – 1/1/2017
DCP – 12.1.10 New Customer Due Diligence	SOP	Original – 1/1/2017
DCP – 12.1.20 New Customer Communications	SOP	Original – 1/1/2017
DCP – 12.2.0 Order Monitoring Program	Policy	Original – 1/1/2017
DCP SOP – 12.2.10 OMP Methodology	SOP	Original – 1/1/2017
DCP SOP – 12.2.11 Product Family Risk Assessment	SOP	Original – 1/1/2017
DCP SOP – 12.2.12 Customer Peer Group Maintenance	SOP	Original – 1/1/2017
DCP SOP – 12.2.20 Identifying and Reporting Suspicious Orders	SOP	Original – 1/1/2017
DCP SOP – 12.2.30 Consumption Reviews	SOP	Original – 1/1/2017
DCP SOP – 12.2.40 OMP Annual Review	SOP	Original – 1/1/2017
DCP SOP – 12.2.50 Annual Audit	SOP	Original – 1/1/2017
DCP – 12.3.0 Ongoing Monitoring Policy	Policy	Original – 1/1/2017
DCP SOP – 12.3.10 Ongoing Monitoring Activities	SOP	Original – 1/1/2017
DCP SOP – 12.3.11 Targeted Pharmacy Visits	SOP	Original – 1/1/2017

¹⁴⁸² See C. Zimmerman memorandum to E. Hazewski, *et al.*, *RVP Talking Points*, 1 (Jan. 19, 2009), ABCMDL00000169.

¹⁴⁸³ See AmerisourceBergen Diversion Control Program Policies & Procedures, ABCMDL00003367 to ABCMDL00003429; D. Mays email to C. Conneely, *et al.*, *Diversion Control Policies* (Jan. 15, 2015) (Caroline Conneely was with FTI), ABCMDL00251385 to ABCMDL00251406.

Document	Policy or SOP	History & Comments
DCP SOP – 12.3.12 – Due Diligence Documentation	SOP	Original – 1/1/2017
DCP SOP – 12.3.20 Communicating Adverse Customer Actions	SOP	Original – 1/1/2017
CSRA 2.12 DCP - Order Monitoring Program	Policy	Original – 12/1/2005 Revised – 6/17/2013 (Document changed to reflect the change in operating systems as well as changes to address additional scrutiny from regulatory/enforcement agencies.)
CSRA 2.25 DCP – Retail Pharmacy Targeted Visits	Policy	Original – 10/1/2008 Revised – 6/17/2013 (Minor changes without changing the substance of policy)
CSRA 2.26 DCP - DEA Daily Reporting	Policy	Original – 10/21/2008
CSRA 2.30 DCP - Suspending CS Shipments to Customers	Policy	Original – 12/5/2013 (Marked “Draft”)
CSRA 3.4 DCP - Customer Account Due Diligence	Policy	Original – 5/8/2007 Revised – 2/13/2013
CSRA 3.5 DCP - Customer Due Diligence Documentation	Policy	Original – 5/10/2013 (This is a new policy)
CSRA 3.9 DCP - GNPPN Accounts Terminated from PBMs	Policy	Original – 7/1/2011

Figure 3: West Virginia Suspicious Order Reports Submitted vs. Dosage Units Shipped¹⁴⁸⁴

Suspicious Order Reports Submitted by AmerisourceBergen to the DEA											
2006	2007*	2008	2009	2010	2011	2012	2013	2014	2015	2016	2017
Number (in Millions) of Oxycodone and Hydrocodone Doses Shipped to West Virginia											

* AmerisourceBergen began to report and block suspicious orders in July 2007; thus, the number of suspicious orders reported in 2007 represents a partial year.

¹⁴⁸⁴ See W. Va. Red Flags Report at 252.

Figure 4: Summit & Cuyahoga Suspicious Orders Reports Submitted¹⁴⁸⁵

County	Year(s)	Number of Orders Reported	
Summit	2007 to 2009		
Summit	2010		
Summit	2011		
Summit	2012		
Summit	2013		
Summit	2014-2018		
Cuyahoga	2007-2008		
Cuyahoga	2009		
Cuyahoga	2010		
Cuyahoga	2011		
Cuyahoga	2012		
Cuyahoga	2013		
Cuyahoga	2014		
Cuyahoga	2015		
Cuyahoga	2016-2018		

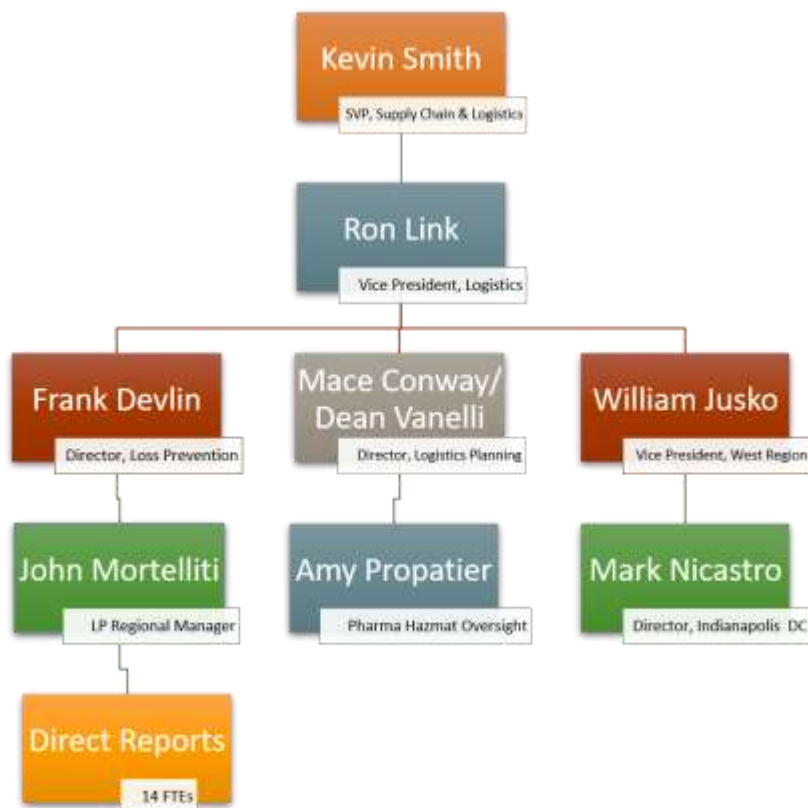
¹⁴⁸⁵ See Spreadsheets ABDCMDL00383974, ABDCMDL00379674, ABDCMDL00383973, ABDCMDL00379673.

Appendix F: CVS Key Facts and Figures

Figure 1: CVS Entity Structure



Figure 2: CVS Logistics Organization



Appendix G: Walgreens Key Facts and Figures

Figure 1: Handling Suspicious Drug Orders

Reproduced below is the text of the Handling Suspicious Drug Orders “policy.”¹⁴⁸⁶

Handling Suspicious Drug Orders

The Logistics and Planning Department sends the Suspicious Control Drug Orders report to all distribution centers. The report lists controlled drug orders that may be:

- Of unusual size for a store in its category
- Of unusual frequency for a store in its category
- Deviating from a normal pattern for a store in Its category

The distribution center must file all reports for five years.

Handling Suspicious Orders and Loss of Controlled Drugs

Policy

Distribution centers must file all Suspicious Control Drug Orders reports for five years. The Administration manager must complete the Report of Theft or Loss of Controlled Substances (DEA Form_106) when any of the following circumstances occur:

- A theft of controlled drugs, no matter how small. (Also file a police report.)
- A substantial loss (a full case or more, an entire repack, or a large dollar amount) of controlled drugs.
- All in-transit losses or thefts, as described above. If store personnel have already signed for the merchandise the pharmacy supervisor is responsible for completing the DEA form

The DEA form must be sent to the DEA, several Walgreens departments, and the local law enforcement agency if the loss was due to theft.

Note: Use U.S. certified mail when sending Form 106 to the DEA. Attach the receipt to the distribution center copy.

¹⁴⁸⁶ See Walgreens, *Handling Suspicious Orders* (Feb. 15, 2005) (emphasis in the original), WAGFLDEA00001854 and WAGFLDEA00001855.

Figure 2: Review of CSR History¹⁴⁸⁷

Phase	Key Points – Scope	Key Points – Identify/Flag selected orders	Key Points – Reduce order quantity for subset of flagged orders
1	<ul style="list-style-type: none"> Deployed in August 2009 Reviews WAG DC orders only Review all Controlled Drug and PSE orders 	<ul style="list-style-type: none"> Flags order based on by drug by store historical sales patterns, i.e. Tolerance threshold or order Frequency 	<ul style="list-style-type: none"> No order reductions in Phase 1
2	<ul style="list-style-type: none"> Deployed in September 2010 No change to scope 	<ul style="list-style-type: none"> No change to Order Identification logic 	<ul style="list-style-type: none"> Automatic reductions to orders that exceed Tolerance threshold
3	<ul style="list-style-type: none"> Deployed in June 2012 Relates vendor orders placed within 48 hours for same drug 	<ul style="list-style-type: none"> Review and refinement of Tolerance/Frequency thresholds. 	<ul style="list-style-type: none"> No change in automatic order reduction logic
4	<ul style="list-style-type: none"> Deployed in August 2012 Incorporates all Vendor orders and partial fills making them eligible for flagging and order reduction. 	<ul style="list-style-type: none"> No change to Order Identification logic 	<ul style="list-style-type: none"> No change in automatic order reduction logic
5	<ul style="list-style-type: none"> Deployed in November 2012 Adds a Ceiling which limits the cumulative receipts of an item by a store 	<ul style="list-style-type: none"> Additional flags created for orders that are in excess of Ceiling for item/store combination Remove Frequency threshold 	<ul style="list-style-type: none"> Automatic reductions to orders that exceed either Tolerance <u>or</u> Ceiling threshold

¹⁴⁸⁷ See Walgreens Presentation, *Controlled Substance Ordering – Evolution of Controlled Substances Ordering Process*, 3 (Oct. 11, 2012) (the system later came to be known as the Controlled Substance Order Monitoring and Prevention System), WAGMDL00667936 at WAGMDL00667938.

Appendix H: Mallinckrodt Key Facts and Figures

Figure 1: Controlled Substances Compliance SOPs

Document Title	Document Number	REVISION DATES						
		2008	2009	2010	2011	2012	2013	2015-18
Identification and Review of Peculiar Orders (precursors to C/S Comp. 3.0) ¹⁴⁸⁸	Draft #1	undated						
	Draft #2	5/13/08						
	Draft #3	6/12/08						
	Draft #4	7/08/08						
	Draft #4	7/15/08						
Identification and Review of Peculiar Orders ¹⁴⁸⁹	C/S Comp 3.0			10/29/10	1/04/11 3/28/11 8/08/11			
Modifications to Procedures for Identification and Review of Unusual Orders ¹⁴⁹⁰					12/08/11			
Identification, Investigation, and Reports of Controlled Substances Suspicious Orders ¹⁴⁹¹	HZQS					9/20/12 10/18/12 11/01/12	3/05/13	08/17/15
New Customer Acct. Set-up & Existing Acct. Ongoing Review ¹⁴⁹²	C/S Comp. 2.0	6/09/08 (precursor)	7/01/09 11/04/09	3/02/10	1/04/11 3/28/11 8/08/11			
Customer Audit Program ¹⁴⁹³	C/S Comp. 4.0				1/04/11 3/28/11 8/08/11			
Identification & Review of	C/S Comp. 5.0				8/08/11			

¹⁴⁸⁸ See MNK-T1_0000273894, MNK-T1_0000268911, MNK-T1_0000419993, MNK-T1_0000296382, MNK-T1_0000263965.

¹⁴⁸⁹ See MNK-T1_0000264260, MNK-T1_0000264275, MNK-T1_0000264200, MNK-T1_0000259166.

¹⁴⁹⁰ See MNK-T1_0000571916.

¹⁴⁹¹ See MNK-T1_0007728766, MNK-T1_0007476261, MNK-T1_0005620500, MNK-T1_0007732477, MNK-T1_0005621914.

¹⁴⁹² See MNK-T1_0000264053, MNK-T1_0000264265, MNK-T1_0000264231, MNK-T1_0000264270, MNK-T1_0000264279, MNK-T1_0000264209, MNK-T1_0000259157.

¹⁴⁹³ See MNK-T1_0000264214, MNK-T1_0000264205, MNK-T1_0000259162.

Suspicious Customer Accts. ¹⁴⁹⁴								
Chargeback Restricting a Pharmacy ¹⁴⁹⁵	HZQS						9/05/13	8/5/17 (draft) 8/14/18 (SpecGx)
Reinstating a Chargeback Restricted Pharmacy ¹⁴⁹⁶						7/16/12	9/05/13	7/28/17 (draft)

The procedures in **blue** represent the keystone documents of Mallinckrodt's suspicious order monitoring program.

Figure 2: "Peculiar Orders" Definition Changes

Definition	Document Title	Doc. Number	Date
"A controlled substance order that meets an internal established criterion that will not be shipped pending further review by DEA Compliance."	Identification and Review of Peculiar Orders (precursors to C/S Comp. 3.0)	Draft #1	undated ¹⁴⁹⁷
(Same as Draft #1)		Draft #2	5/13/08 ¹⁴⁹⁸
"A controlled substance order that meets an internal established criterion that will be placed on hold pending further review by DEA Compliance."		Draft #3	6/02/08 ¹⁴⁹⁹
(Same as Draft #3)		Draft #4	7/08/08 ¹⁵⁰⁰
(Same as Draft #3)		Draft #4	7/15/08 ¹⁵⁰¹
"Controlled substance order that meets internal established criteria of 3X the average amount of product ordered during the previous 12 months by DEA reporting class."	Identification and Review of Peculiar Orders	C/S Comp 3.0	10/29/10 ¹⁵⁰²
(Same as 10/29/10 version)	Identification and Review of Peculiar Orders	C/S Comp 3.0	1/04/11 ¹⁵⁰³
"Mallinckrodt direct customer controlled substance order that meets internal established criteria of 3X the average amount of product ordered during the previous 12 months by DEA reporting class."	Identification and Review of Peculiar Orders	C/S Comp 3.0	3/28/11 ¹⁵⁰⁴
(Same as 3/28/11)	Identification and Review of	C/S Comp 3.0	8/08/11 ¹⁵⁰⁵

¹⁴⁹⁴ See MNK-T1_0000259153.

¹⁴⁹⁵ See MNK-T1_0000511225; MNK-T1_0007732565; MNK-T1_0004155830.

¹⁴⁹⁶ See MNK-T1_0007732447; MNK-T1_0007732532; MNK-T1_0007732621

¹⁴⁹⁷ MNK-T1_0000273894

¹⁴⁹⁸ MNK-T1_0000268911.

¹⁴⁹⁹ MNK-T1_0000419993.

¹⁵⁰⁰ MNK-T1_0000296382.

¹⁵⁰¹ MNK-T1_0000263965.

¹⁵⁰² MNK-T1_0000264260.

¹⁵⁰³ MNK-T1_0000264275.

¹⁵⁰⁴ MNK-T1_0000264200.

Definition	Document Title	Doc. Number	Date
(Not Mentioned)	Peculiar Orders		12/8/11 ¹⁵⁰⁶
	Modifications to Procedures for Identification and Review of Unusual Orders		
“A standard algorithm with respect to volume which sets a monthly limit at: (a) 2.5X the average number of orders of a product during the previous 18 months by the customer and (b) 2.5X the average volume of product ordered during the previous 18 months by the customer.”	Identification, Investigation, and Reports of Controlled Substances Suspicious Orders	HZQS	11/01/12 ¹⁵⁰⁷

¹⁵⁰⁵MNK-T1_0000259166.

¹⁵⁰⁶MNK-T1_0000571916.

¹⁵⁰⁷MNK-T1_0005620500

Appendix I: List of Materials Considered

A. Defendant Production Documents

CAH_MDL_PRIORPROD_DEA07_01185382	WAGMDL00325170
CAH_MDL_PRIORPROD_DEA12_00011059	WAGMDL00325129
HDS_MDL_00002032	WAGMDL00667936
Acquired_Actavis_00441354	WAGMDL00658227
CAH_MDL2804_01431074	WAGMDL00246016
WAGMDL00490963	WAGMDL00010925
WAGMDL00493697	WAGMDL00010926
WAGMDL00493694	WAGMDL00010927
WAGMDL00387635	WAGMDL00010928
WAGMDL00387641	WAGMDL00037093
WAGMDL00709395	WAGMDL00037094
WAGMDL00749381	WAGMDL00095316
WAGMDL00010887	WAGMDL00095317
WAGMDL00751821	ALLERGAN_MDL_03755273
WAGMDL00751871	MCKMDL00478906
WAGMDL00777158	MCKMDL00478910
WAGMDL00414048	MNK-T1_0001454856
WAGMDL00303305	WAGFLDEA00000846
WAGMDL00060486	WAGFLDEA00000852
WAGMDL00400358	WAGMDL00006645
WAGMDL00400360	WAGMDL00021425
WAGMDL00101723	WAGMDL00035669
WAGMDL00624503	WAGMDL00044765
WAGMDL00659801	WAGMDL00077015
WAGMDL00660331	WAGMDL00077016
WAGMDL00659270	WAGMDL00102390
WAGMDL00674280	WAGMDL00107267
WAGMDL00574824	WAGMDL00107557
WAGMDL00358471	WAGMDL00113808
WAGMDL00400357	WAGMDL00119539
WAGMDL00700240	WAGMDL00119542
WAGMDL00709508	WAGMDL00183798
WAGMDL00709510	WAGMDL00237698
WAGMDL00477975	WAGMDL00245867
WAGMDL00107532	WAGMDL00254645
WAGMDL00415348	WAGMDL00254649
WAGMDL00757193	WAGMDL00289068
WAGMDL00757759	WAGMDL00302174
WAGMDL00757762	WAGMDL00317093
WAGMDL00757788	WAGMDL00319129
WAGFLDEA00001854	WAGMDL00325368
WAGFLDEA00000027	WAGMDL00379420
WAGFLDEA00000028	WAGMDL00386743
WAGFLDEA00000117	WAGMDL00387625

WAGMDL00387629
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WAGMDL00387631
WAGMDL00387632
WAGMDL00387633
WAGMDL00387638
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WAGMDL00394499
WAGMDL00395957
WAGMDL00396133
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CAH_MDL2804_00012245	MNK-T1_0000559412
CAH_MDL2804_00012249	MNK-T1_0000559532
MNK-T1_0000259166	MNK-T1_0000559581
MNK-T1_0000263874	MNK-T1_0000559994
MNK-T1_0000263875	MNK-T1_0000560227
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MNK-T1_0000264260	MNK-T1_0000561015
MNK-T1_0000264275	MNK-T1_0000562520
MNK-T1_0000264292	MNK-T1_0000563394
MNK-T1_0000267896	MNK-T1_0000563696

MNK-T1_0000571916	ABDCMDL00396853
MNK-T1_0004888229	ABDCMDL00395454
MNK-T1_0007146630	ABDCMDL00396658
MNK-T1_0007728295	ABDCMDL00396567
CAH_MDL2804_01457737	ABDCMDL00396568
CAH_MDL_PRIORPROD_HOUSE_0000264	ABDCMDL00396656
CAH_MDL2804_02953369	MNK-T1_0007728766
MNK-T1_0000259153	MNK-T1_0007732447
MNK-T1_0000259157	MNK-T1_0007732477
MNK-T1_0000259162	MNK-T1_0007732532
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MNK-T1_0000264209	MNK-T1_0000269039
MNK-T1_0000264214	MNK-T1_0000280607
MNK-T1_0000264231	MNK-T1_0000280835
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MNK-T1_0000511227	MNK-T1_0000472004
MNK-T1_0000511246	MNK-T1_0000473333
MNK-T1_0002357607	MNK-T1_0000383311
MNK-T1_0004155827	MNK-T1_0000384265
MNK-T1_0004155830	MNK-T1_0004155830
MNK-T1_0004155833	MNK-T1_0000368390
MNK-T1_0005620500	MNK-T1_0000372333
MNK-T1_0005621914	MNK-T1_0000562682
MNK-T1_0007476261	MNK-T1_0000264240
MCKMDL00647803	MNK-T1_0000296081
ENDO-OPIOID_MDL-02776076	MNK-T1_0000307203
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ABDCMDL00170149	MNK-T1_0000565624
PPLPC039000161600	MNK-T1_0007097723
ABDCMDL00331149	MNK-T1_0000289708
ABDCMDL00331154	MNK-T1_0000312043
CAH_MDL2804_02509732	MNK-T1_0000269410
CAH_MDL2804_03262274	MNK-T1_0007063163
CAH_MDL2804_01287246	MNK-T1_0005032855
MNK-T1_0000268860	MNK-T1_0006029397
CVS-MDLT1-000010268	MNK-T1_0002336044
MCKMDL00407451	MNK-T1_0006031387
ENDO-OPIOID_MDL-03660023	MNK-T1_0007703635
ABDCMDL00354768	MNK-T1_0002158106
ABDCMDL00354771	MNK-T1_0007703718
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ABDCMDL00396256	MNK-T1_0001806022
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ABDCMDL00396821	MNK-T1_0002734994

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MNK-T1_0000496062	CVS-MDLT1-000124108
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CVS-MDLT1-000007490	CVS-MDLT1-000124124
CVS-MDLT1-000101092	CVS-MDLT1-000124142
CVS-MDLT1-000101174	CVS-MDLT1-00012414
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CVS-MDLT1-000003028	
CVS-MDLT1-000010552	
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CVS-MDLT1-000123843	
CVS-MDLT1-000123857	
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CVS-MDLT1-000123912	
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- Laura Merton LinkedIn Profile, <https://www.linkedin.com/in/lauramerten/> (last accessed Feb. 12, 2019)
- Matthew D'Ambrosio LinkedIn Profile, <https://www.linkedin.com/in/mattdambrosio/> (last accessed Feb. 12, 2019).

F. Defendant Discovery Responses; MDL No. 2804 *IN RE: National Prescription Opiate Litigation*

- AmerisourceBergen's Second Supplemental Objections and Responses to Plaintiffs' Combined Discovery Requests (Mar. 4, 2019)

- Cardinal Health Inc.'s Third Supplemental Objections and Responses to Plaintiffs' First Combined Discovery Requests (Mar. 4, 2019)
- CVS RX Services, Inc.'s and CVS Indiana L.L.C.'s Objections and Responses to Plaintiffs' (First) Combined Discovery Requests to National Retail Pharmacy Defendants (Nov. 30, 2018)
- McKesson Corporation's Second Supplemental Objections and Responses to Plaintiffs' Combined Discovery Requests (Mar. 4, 2019)
- McKesson Corporation's Third Supplemental Objections and Responses to Plaintiffs' Combined Discovery Requests Nos. 2 and 3 (March 29, 2019)
- Walgreen Co. and Walgreen Eastern Co.'s Second Amended Objections and Responses to Plaintiffs' First Set of Interrogatories (Mar. 4, 2019)
- Mallinckrodt's Supplemental Responses and Objections to Interrogatory Nos. 1, 5, 7, 8, 9, 16, 21, 22, 23, 27, 30, 31, 32, 33, and 35 (Jan. 30, 2019)
- Mallinckrodt's Supplemental Responses and Objections to Interrogatory Nos. 1-5, 7-9, 11-13, 15-16, 18-19, 21-22, 25-26, and 29-35 (Mar. 5, 2019)

G. Corporate Witness Depositions

- Deposition of Nathan Elkins + Exhibits (11/14/2018) (AmerisourceBergen)
- Deposition of Edward Hazewski + Exhibits (10/25/2018) (AmerisourceBergen)
- Deposition of David May + Exhibits (8/4/2018) (AmerisourceBergen)
- Deposition of Steve Mays + Exhibits (10/24/2018) (AmerisourceBergen)
- Deposition of Chris Zimmerman + Exhibits (8/3/2018) (AmerisourceBergen)
- Deposition of Steve Mays - Volume II + Exhibits (2/8/2019) (AmerisourceBergen)
- Deposition of Marcelino Guerreiro + Exhibits (4/3/2019) (AmerisourceBergen)
- Deposition of Mark Hartman + Exhibits (11/15/2018) (Cardinal)
- Deposition of Kim Howenstein + Exhibits (1/10/2019) (Cardinal)
- Deposition of Stephen Reardon + Exhibits (11/30/2018) (Cardinal)
- Deposition of Chris Lancot + Exhibits (10/10/2018) (Cardinal)
- Deposition of Steve Morse + Exhibits (12/13/2018) (Cardinal)
- Deposition of Christopher Forst + Exhibits (1/22/2019) (Cardinal)
- Deposition of Jennifer Norris + Exhibits (8/7/2018) (Cardinal)
- Deposition of Eric Brantley + Exhibits (11/27/2018) (Cardinal)
- Deposition of Nick Rausch + Exhibits (11/16/2018) (Cardinal)
- Deposition of Steve Lawrence + Exhibits (1/4/2019) (Cardinal)
- Deposition of Kelly Baker + Exhibits (1/24/2019) (CVS)
- Deposition of Aaron Burtner + Exhibits (1/17/2019) (CVS)
- Deposition of Frank Devlin + Exhibits (1/10/2019) (CVS)
- Deposition of Terrance Dugger + Exhibits (1/23/2019) (CVS)
- Deposition of Shauna Helfrich + Exhibits (1/10/2019) (CVS)
- Deposition of Pam Hinkle + Exhibits (1/24/2019) (CVS)
- Deposition of Sherri Hinkle + Exhibits (1/25/2019) (CVS)
- Deposition of Ronald Link + Exhibits (12/11/2018) (CVS)
- Deposition of Gary Milikan + Exhibits (1/11/2019) (CVS)
- Deposition of John Mortelliti + Exhibits (1/23/2019) (CVS)
- Deposition of Mark Nicastro + Exhibits (12/6/2018) (CVS)
- Deposition of Amy Propatier + Exhibits (11/29/2018) (CVS)
- Deposition of Craig Shiavo + Exhibits (1/17/2019) (CVS)
- Deposition of Dean Vanelli + Exhibits (1/16/2019) (CVS)

- Deposition of Mark Vernazza + Exhibits (11/20/2018) (CVS)
- Deposition of Ellen Wilson + Exhibits (1/24/2019) (CVS)
- Deposition of Dave Gustin + Exhibits (8/17/2018) (McKesson)
- Deposition of Michael Oriente + Exhibits (7/19/2018) (McKesson)
- Deposition of Blaine Snider + Exhibits (11/8/2018) (McKesson)
- Deposition of Gary Hilliard + Exhibits (1/10/2019) (McKesson)
- Deposition of Donald Walker + Exhibits (1/10/2019) (McKesson)
- Deposition of Nate Hartle (30(b)(6)) + Exhibits (7/31/2018) (McKesson)
- Deposition of Nate Hartle + Exhibits (8/1/2018) (McKesson)
- Deposition of William De Gutierrez-Mahoney + Exhibits (11/28/2018) (McKesson)
- Deposition of Gene Cavacini + Exhibits (1/25/2019) (McKesson)
- Deposition of Gary Boggs + Exhibits (7/19/2018) (McKesson)
- Deposition of Gary Boggs + Exhibits (1/17/2019) (McKesson)
- Deposition of Tracy Jonas + Exhibits (11/15/2018) (McKesson)
- Deposition of Tom McDonald + Exhibits (12/07/2018) (McKesson)
- Deposition of Micheal Bishop + Exhibits (1/09/2019) (McKesson)
- Deposition of Stephen Bamberg + Exhibits (12/14/2018) (Walgreens)
- Deposition of Wayne Bancroft + Exhibits (1/10/2019) (Walgreens)
- Deposition of Edward Bratton + Exhibits (11/30/2018) (Walgreens)
- Deposition of Edward Bratton (30 (b)(6)) + Exhibits (12/16/2018) (Walgreens)
- Errata For Deposition of Edward Bratton (30 (b)(6)) (12/16/2018) (Walgreens)
- Deposition of Christopher Dymon + Exhibits (1/25/2019) (Walgreens)
- Deposition of Barbara Martin + Exhibits (1/25/2019) (Walgreens)
- Deposition of John Merritello + Exhibits (1/18/2019) (Walgreens)
- Deposition of Steve Mills + Exhibits (11/8/2018) (Walgreens)
- Deposition of Denman Murray Jr. + Exhibits (1/15/2019) (Walgreens)
- Deposition of Natasha Polster + Exhibits (1/23/2019) (Walgreens)
- Deposition of Eric Stahmann + Exhibits (10/16/2018) (Walgreens)
- Deposition of Rex Swords + Exhibits (12/21/2018) (Walgreens)
- Deposition of Deborah Bish+ Exhibits (2/1/2019) (Walgreens)
- Deposition of Jennifer Diebert + Exhibits (1/24/2019) (Walgreens)
- Deposition of John Adams + Exhibits (1/31/2019) (Mallinckrodt)
- Deposition of Steven Becker + Exhibits (12/19/2018) (Mallinckrodt)
- Deposition of Victor Borelli + Exhibits (11/29/2018) (Mallinckrodt)
- Deposition of Ginger Collier + Exhibits (1/08/2019) (Mallinckrodt)
- Deposition of John Gillies + Exhibits (2/07/2019) (Mallinckrodt)
- Deposition of Karen Harper + Exhibits (1/15/2019) (Mallinckrodt)
- Deposition of Kate Neely (Muhlenkamp) + Exhibits (1/08/2019) (Mallinckrodt)
- Deposition of William Ratliff + Exhibits (12/19/2018) (Mallinckrodt)
- Deposition of James Rausch + Exhibits (11/16/2018) (Mallinckrodt)
- Deposition of Tiffany Rowley-Kilper + Exhibits (2/09/2019) (Mallinckrodt)
- Deposition of Cathy Stewart + Exhibits (12/11/2018) (Mallinckrodt)
- Deposition of Hugh O'Neill + Exhibits (3/13/2019) (Mallinckrodt)

H. Third-Party Witness Depositions

- Deposition of Kyle Wright – Volume I + Exhibits (2/28/2019) (DEA)
- Deposition of Kyle Wright – Volume II + Exhibits (3/4/2019) (DEA)

- Deposition of Demetra Ashley + Exhibits (3/15/2019) (DEA)

I. Other Non-Publicly-Available Materials

- Cardinal Health Organization Chart 2012-2015 (P1.4592)
- Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 218-221, 1944-1947 (Oxycodone and Hydrocodone Distribution to CVS Pharmacy #03322 by Cardinal Health and CVS; Opioid Shipments to AR7531418 by Distributor; Ohio CVS Stores, LLC, AR7531418 Total Dosage Units and Total MME by Drug Family)
- Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 1416-1419, 3216-3219 (Oxycodone and Hydrocodone Distribution to CVS Pharmacy #04800 by Cardinal Health and CVS; Opioid Shipments to BR0287234 by Distributor; Ohio CVS Stores, LLC, BR0287234 Total Dosage Units and Total MME by Drug Family)
- Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 51, 136 (Total Dosage of Oxycodone and Hydrocodone Shipped by Cardinal Health to Cuyahoga County, Ohio (Jan. 1996 to April 2018))
- Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 93, 178 (Total Dosage of Oxycodone and Hydrocodone Shipped by Cardinal Health to Summit County, Ohio (Jan. 1996 to April 2018))
- Appendix 9 to Expert Report of Craig J. McCann, Ph.D., CFA dated March 25, 2019 at 318 (Oxycodone Distribution to Euclid Family Pharmacy)

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WWW.WHITELAWCOMPLIANCE.COM

SWHITELAW@WHITELAWCOMPLIANCE.COM

DR. SETH B. WHITELAW

PROFESSIONAL SUMMARY

Dr. Whitelaw has more than 25 years of industry experience in the life sciences and healthcare sectors, as an attorney, compliance officer and consultant. His career has focused on food and drug law and corporate governance, as well as designing and running compliance programs within medical devices, pharmaceutical sales and marketing, and pharmaceutical R&D. He is a licensed food and drug attorney, with a doctorate in Health Law. His forte is designing, building and running life science compliance programs from a "blank sheet of paper."

LICENSES & INTERNSHIPS

Licensed to Practice Law in the Commonwealths of Pennsylvania (2004) and Virginia (1988)

Food and Drug Law Institute Fellowship (1988)

Internship with U.S. Food and Drug Administration, Office of Chief Counsel (1988-1989)

Internships with Grocery Manufacturers Association (GMA), Washington, D.C.

EXPERIENCE

WHITELAW COMPLIANCE GROUP, LLC., PHILADELPHIA, PA **President & CEO, April 2015 – Present**

Focused exclusively on small to medium-sized FDA-regulated companies, the Whitelaw Compliance Group provides practical, pragmatic compliance and integrity services that are tailored to each regulated company to help them grow and achieve sustainable integrity.

Responsible for designing, developing and implementing a med tech tailored compliance program to address compliance risks, including FCPA and UK Bribery Act issues.

POLICY & MEDICINE COMPLIANCE UPDATE, COLUMBIA, MD **Editor, October 2015 – Present**

Formerly Life Science Compliance Update. Oversees the editorial content, assembly and monthly publication providing comprehensive, up-to-date compliance information for pharmaceutical, biotechnology, and device manufacturers. Writes articles on emerging life sciences issues for the publication.

MITCHELL HAMLINE SCHOOL OF LAW, ST. PAUL, MN **Senior Fellow and Adjunct Professor, Life Sciences Compliance, September 2016 – Present**

Oversaw, designed and taught Legal Compliance Essentials for Drug, Device and Biotech Companies (J-Term 2017)

Co-teaching Health Care Compliance Skills (Fall 2017 & Spring-Fall 2018)

MISONIX, INC., FARMINGDALE, NY
Interim Chief Compliance Officer, December 2016 – June 2017

Interim Chief Compliance Officer for Misonix, Inc., which specializes in the development and commercialization of ultrasonic surgical devices for neurosurgical, spinal, advanced wound care, and general surgery procedures. Responsible for the day-to-day implementation and operation of the Compliance Program including compliance efforts involving interactions with health care professionals and anti-bribery/anti-corruption.

- Delivered over \$165K in sales, from Chinese distributors as a result of overseeing and managing enhanced third-party due diligence and contracting process.

DELOITTE & TOUCHE LLP., PHILADELPHIA, PA
Director, October 2011 – April 2015

Led the Advisory Practice's transparency team assisting U.S. and other global medical device and pharmaceutical clients in developing effective processes and operating approaches to meet both U.S. Sunshine Act requirements, as well as other global requirements (e.g., France, Japan and EFPIA).

- Consistently delivered more than \$1M in sales each year.

Advised various multinational clients on structuring a global compliance function including work plan prioritization.

Assisted a client with medical device, pharmaceutical, and consumer products units to develop a streamlined and strategic Medical Affairs department to support its globally growing business.

Conducted multiple internal audits for clients in both the R&D and third-party oversight areas working with global teams.

Served as Editor-in-Chief and contributing author for Deloitte's @Regulatory bulletins from 2013-2015

GLAXOSMITHKLINE, PHILADELPHIA, PA
Compliance Officer, Global R&D, January 2001 – October 2011

Successfully designed, developed, implemented and led the corporate compliance infrastructure, including integrating internal audit with compliance, for GSK's global R&D operations where none had existed previously.

Founded and supported the compliance infrastructure for GSK's new R&D China site in Shanghai.

Provided compliance oversight and support to sites in U.S, U.K. China, Italy, Spain, France and Croatia with small (9) central staff on a broad range of topics including conflicts of interest, anti-kickback, FCPA, false claims, use of human biological samples, transparency, etc.

Created and implemented policies, systems and processes ahead of industry practice to reduce the risk from perceived improper influence with healthcare professionals, especially in countries with national health insurance programs.

Successfully negotiated with various regulatory authorities to resolve compliance issues.

Led compliance efforts surrounding GSK's voluntary disclosure of research payments to healthcare professionals and healthcare institutions (e.g., transparency).

Helped lead R&D's efforts to prepare for impending Corporate Integrity Agreement

SMITHKLINE BEECHAM PHARMACEUTICALS, PHILADELPHIA, PA

Legal Compliance Officer, January 1997 – January 2001

Successfully designed and implemented the corporate compliance infrastructure for the U.S. and Canadian commercial operations where none had existed previously and co-led the integration of the departments during the Glaxo Wellcome/SmithKline merger.

Created and implemented policies, systems and processes ahead of industry practice to reduce the risk from perceived improper influence with healthcare professionals (e.g., banning gifts).

Successfully help lead the efforts to enhance SmithKline's sample accountability (PDMA) program.

C.R. BARD, INC., MURRAY HILL, NJ

Senior Attorney & Compliance Coordinator, January 1991 - January 1997

Created and implemented Bard's original corporate medical device compliance program to meet the requirements of the Federal Sentencing Guidelines and Bard's Plea Agreement with the U.S. Department of Justice and served as Bard's first Compliance Officer post settlement.

Successfully directed and managed Bard's company-wide document production efforts for the U.S. v. C.R. Bard, Inc. litigation resulting in the production of over 750,000 responsive pages. Worked directly with both the AUSA's Office in Boston as well as FDA's Office of Criminal Investigation.

Oversaw and updated Bard's records retention program.

Created, managed and implemented a Legal Audit Program to provide the Corporation with a concrete evaluation of its overall compliance with both the federal FDA regulatory scheme (e.g., 501(k) compliance, Quality System requirements) and its internal policies. This program was successfully integrated with Bard's already established internal and quality auditing programs.

Provided legal counsel on various medical device regulatory matters including those issues involving FDA, EPA and OSHA (e.g., custom devices, consumer preference testing, medical device reports, FDA-483 and Warning Letter responses)

FD Inc., Washington, D.C.

Head of Sales and Marketing, March 1990- January 1991

Sales and marketing of food and drug statutory, administrative and regulatory materials on compact disk, with direct responsibility for developing and implementing both short and long-term marketing strategies for the company.

Fox, Bennett & Turner., Washington, D.C.

Associate, May 1989- March 1990

Compliance counseling and opinion drafting on food, drug and environmental issues, particularly safety and risk assessment.

EDUCATION

WIDENER UNIVERSITY SCHOOL OF LAW, WILMINGTON, DE
2011 – S.J.D., Health Law

GEORGE WASHINGTON UNIVERSITY LAW SCHOOL, WASHINGTON, D.C.
1989- LL.M., Administrative Law

WASHINGTON & LEE UNIVERSITY, SCHOOL OF LAW, LEXINGTON, VA
1988 – J.D.

BOWDOIN COLLEGE, BRUNSWICK MAINE
1985- A.B., History (*Cum laude*)

Publications List

PUBLICATIONS

- Whitelaw, Seth; Fiorentino, Nicodemo; and O'Leary, Jennifer "Drug Pricing—The Next Compliance Waterloo," Mitchell Hamline Law Review: 2018 Vol. 44: Iss. 4, Article 2. Available at: <https://open.mitchellhamline.edu/mhlr/vol44/iss4/2>.
- Schroeder, Whitelaw, Makosch, Adapt or Perish -Can Stem Cell Therapies Achieve Their Potential for Delivering Optimal, Cost-Effective Clinical Outcomes in an Evolving Regulatory Framework?, Life Science Compliance Update Special Supplement (Aug. 2018)
- Whitelaw, et al., The Day After Tomorrow - The Drug Pricing Chorus Grows Louder, 4.4 Life Science Compliance Update 1 (Apr. 2018)
- “Missing the Market: Government Standards Are Undermining Compliance Efforts in Smaller Life Science Companies,” Attorney at Law Magazine, Minnesota Ed. (Mar. 2018)
- Whitelaw, et al., A Bright Future or Unfulfilled Promise – An Update on Biosimilars and Their Prospects for Contributing to Meaningful Cost Reduction, 4.3 Life Science Compliance Update 13 (Mar. 2018).
- “One Purpose to Rule Them All – A Resounding ‘Yes’ According to the District Court in U.S. ex rel. Cairns,” Life Science Compliance Update, Vol. 4.2 (Feb. 2018).
- “On a Collision Course – FDA Clinical Investigator Disclosure and Open Payments,” Life Science Compliance Update, Vol. 2.9 (Sep. 2016).
- “The Board of Directors’ Role in Pharmaceutical Compliance,” Pharmaceutical Compliance Monitor (Dec. 10, 2012).
- Evaluating IRB’s and Their Roles, 16 Food, Drug, Cosmetic and Medical Device Law Digest
- “How Can FDA Improve Its Financial Disclosure Rules for Clinical Investigators in this New Era of Transparency?”, Food and Drug Law Institute Policy Forum (Jun. 2011).
- “Proposition 65 v. Industry: David Against Goliath or a Misled Public Run Amok?,” 44 Food Drug Cosmetic Law Journal 677
- “FDA Publishes The New UDI Regulations – Will You Be Ready?,” Deloitte (Oct. 2013)
http://www.deloitte.com/view/en_US/us/Industries/health-care-providers/9026855ddef61410VgnVCM1000003256f70aRCRD.htm

- “Four Actions You Can Still Take to Begin Sunshine Act Compliance”, Deloitte (Aug. 2013)
http://www.deloitte.com/view/en_US/us/Insights/centers/center-regulatory-strategies/crs-blog/d8f713e7e1c90410VgnVCM2000003356f70aRCRD.htm)
- “Time Crunch - Physician Payments Sunshine Act,” Deloitte (Jun. 2013)
(http://www.deloitte.com/view/en_US/us/Services/audit-enterprise-risk-services/governance-regulatory-risk-strategies/9a0af1b41c08f310VgnVCM1000003256f70aRCRD.htm)
- “The Board of Directors’ Role in Pharmaceutical Compliance,” Pharmaceutical Compliance Monitor (Dec. 10, 2012), <http://www.pharmacompliancemonitor.com/the-board-of-directors-role-in-pharmaceutical-compliance-2/3677/>
- Practicing Avoidance: Navigating Qui Tam and Consent Decrees, Pharmaceutical Compliance Monitor (Jan. 9, 2012), <http://www.pharmacompliancemonitor.com/practicing-avoidance-navigating-qui-tam-consent-decrees/#more-996>.
- How Can FDA Improve Its Financial Disclosure Rules for Clinical Investigators in this New Era of Transparency?, Food and Drug Law Institute Policy Forum (June 2011)

ONLINE CONTENT (EDITOR)

- Policy & Medicine Compliance Update Vol. 5.4 April 2019.
- Policy & Medicine Compliance Update Vol. 5.3 March 2019.
- Policy & Medicine Compliance Update Vol. 5.2 February 2019.
- Policy & Medicine Compliance Update Vol. 5 January 2019.
- Policy & Medicine Compliance Update Vol. 4.12 December 2018.
- Policy & Medicine Compliance Update Vol. 4.11 November 2018.
- Policy & Medicine Compliance Update Vol. 4.10 October 2018.
- Policy & Medicine Compliance Update Vol. 4.9 September 2018.
- Adapt or Perish Can Stem Cell Therapies Achieve Their Potential For Delivering Optimal Cost-Effective Clinical Outcomes In an Evolving Regulatory Framework?, Policy & Medicine Life Science Compliance Special Supplement Vol. 4.8 August 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.8 August 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.7 July 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.6 June 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.5 May 2018.

- Policy & Medicine Life Science Compliance Update Vol. 4.4 April 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.3 March 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.2 February 2018.
- Policy & Medicine Life Science Compliance Update Vol. 4.1 January 2018.
- Policy & Medicine Life Science Compliance Update Vol. 3.12 December 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.11 November 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.10 October 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.9 September 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.8 August 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.7 July 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.6 June 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.5 May 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.4 April 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.3 March 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.2 February 2017.
- Policy & Medicine Life Science Compliance Update Vol. 3.1 January 2017.
- Policy & Medicine Life Science Compliance Update Vol. 2.12 December 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.11 November 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.10 October 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.9 September 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.8 August 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.7 July 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.6 June 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.5 May 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.4 April 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.3 March 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.2 February 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.1 January 2016.
- Policy & Medicine Life Science Compliance Update Vol. 2.0 December 2015.
- Policy & Medicine Life Science Compliance Update Vol. 1.9 November 2015.

- Compliance for the Common Man Policy & Medicine Life Science Compliance Update Vol. 1.5 July 2015.

REPRESENTATIVE SPEAKING ENGAGEMENTS

- Guest lecturer at Temple University, Ursinus College, Medical Devices Section, Food and Drug Law, Rutgers-Camden Law School
- Mitchell Hamline's Health Law Institute Symposium - Hot Topics in Healthcare Compliance (2018)
- CBI 2nd Annual Drug Pricing Transparency Conference (2018)
- CBI Annual Pharmaceutical Compliance Congress (multiple years) Pharmaceutical Regulatory and Compliance Congress (multiple years)
- Mitchell Hamline School of Law, National Speaker Series (Oct. 2016)
- DIA Marketing Pharmaceuticals (2014)
- Sixth National Disclosure Summit (2014)
- Pharmaceutical Regulatory and Compliance Congress (2013)
- Food and Drug Law Institute Advertising and Promotion Conference (2013)
- CBI 9th Annual Pharmaceutical Accounting and Reporting Congress (2013)
- AdvaMed Conference (2012)
- CBI 9th Annual Pharmaceutical Compliance Congress (2012)
- FDLI US-China Food and Drug Law (2011)
- Food and Drug Law Institute Annual Conference (2011)
- ACI 11th National Forum on Fraud & Abuse in Sales & Marketing (2011)
- Widener Law 2d Annual Regulatory Compliance Program (2010)
- Marcus Evans Commercial Compliance for Pharmaceutical & Medical Device Companies (2010)
- CBI Clinical R&D Compliance Forum (2010)
- Drug Information Association 46th Annual Meeting (2010)
- Pharma Compliance Forum (2009 & 2010)
- ACI Medical Affairs Conference (2009)

Prior Testimony

None.